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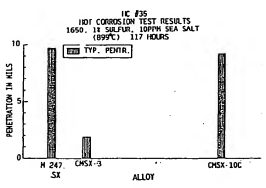
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- (54) Single crystal nickel-based superalloy.
- A nickel-based superalloy comprises the following elements in percent by weight: from about 5.0 to about 7.0 percent rhenium, from about 1.8 to about 4.0 percent chromium, from about 1.5 to about 9.0 percent cobalt, from about 7.0 to about 10.0 percent tantalum, from about 3.5 to about 7.5 percent tungsten, from about 5.0 to about 7.0 percent aluminium, from about 0.1 to about 1.2 percent titanium, from about 0.5 percent columbium, from about 0.25 to about 2.0 percent molybdenum, from about 0 to about 0.15 percent hafnium, and the balance nickel + incidental impurities, the superalloy having a phasial stability number N<sub>V3B</sub> less than about 2.10.



TYP. PENTR. = AVERAGE DEPTH OF CORROSION ATTACK ON PINS IN GROUP

Fig. 1

Jouve, 18, rue Saint-Denis, 75001 PARIS

#### BACKGROUND OF THE INVENTION

#### 1. Field of the Invention

This invention relates to single crystal nickel-based superalloys and, more particularly, single crystal nickel-based superalloys and articles made therefrom for use in advanced gas turbine engines under high stress, high temperature conditions.

#### 2. Description of the Prior Art

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Advances over recent years in the metal temperature and stress capability of single crystal articles have been the result of the continuing development of single crystal superalloys, as well as improvements in casting processes and engine application technology. These single crystal superalloy articles include rotating and stationary turbine blades and vanes found in the hot sections of gas turbine engines. However, gas turbine engine design goals have remained the same during the past decades. These goals include the desire to increase engine operating temperature, rotational speed, thrust-to-weight ratio, fuel efficiency, and engine component durability and reliability.

The basic technology of alloys for the casting of single crystal components is described in U.S. Patent Nos. 3,494,709; 4,116,723 and 4,209,348. Development work resulted in first generation nickel-based superalloys, which were materially improved over those described in the aforementioned patents. However, these first generation nickel-based superalloys contained no rhenium. Examples of such first generation nickel-based superalloys, commercially known as CMSX-2 alloy and CMSX-3 alloy produced by Cannon-Muskegon Corporation, assignee of the present application, are described in U.S. Patent No. 4,582,548. Further development work resulted in second generation nickel-based superalloys hating improved creep strength/creep rate. These second generation nickel-based superalloys have a moderate rhenium content of about 3 weight percent. An example of such a second generation nickel-based superalloy is described in U.S. Patent No. 4,643,782. This patent discloses a superalloy, commercially known as CMSX-4 alloy, having a specific nickel-based composition including a rhenium content in the range of 2.8-3.2 weight percent. The present invention provides the next generation of nickel-based superalloys having higher total refractory element (W+Re+Mo+Ta) content and improved mechanical properties.

Single crystal articles are generally produced having the low-modulus (001) crystallographic orientation parallel to the component dendritic growth pattern or blade stacking fault axis. Face-centered cubic (FCC) superalloy single crystals grown in the (001) direction provide extremely good thermal fatigue resistance relative to conventionally cast articles. Since these single crystal articles have no grain boundaries, alloy design without grain boundary strengtheners, such as carbon, boron and zirconium, is possible. As these elements are alloy melting point depressants, their reduction from an alloy design provides a greater potential for high temperature mechanical strength achievement since more complete gamma prime solution and microstructural homogenization can be achieved relative to directionally solidified (DS) columnar grain and conventionally cast materials. Their reduction also makes possible a higher incipient melting temperature.

These process benefits are not necessarily realized unless a multi-faceted alloy design approach is undertaken. Alloys must be designed to avoid tendency for casting defect formation such as freckles, slivers, spurious grains and recrystallization. Additionally, the alloys must provide an adequate heat treatment window (numeric difference between an alloy's gamma prime solvus and incipient melting point) to allow for nearly complete gamma prime solutioning. At the same time, the alloy compositional balance should be designed to provide an adequate blend of engineering properties necessary for operation in gas turbine engines. Selected properties generally considered important by gas turbine engine designers include: elevated temperature creep-rupture strength, thermo-mechanical fatigue resistance, impact resistance plus hot corrosion and oxidation resistance.

An alloy designer can attempt to improve one or two of these design properties by adjusting the compositional balance of known superalloys. However, it is extremely difficult to improve more than one or two of the design properties without significantly or even severely compromising the remaining properties. The unique superalloy of the present invention provides an excellent blend of the properties necessary for use in producing single crystal articles for operation in gas turbine engine hot sections.

## SUMMARY OF THE INVENTION

This invention relates to a nickel-based superalloy comprising the following elements in percent by weight: from about 5.0 to about 7.0 percent rhenium, from about 1.8 to about 4.0 percent chromium, from about 1.5 to

about 9.0 percent cobalt, from about 7.0 to about 10.0 percent tantalum, from about 3.5 to about 7.5 percent tungsten, from about 5.0 to about 7.0 percent aluminum, from about 0.1 to about 1.2 percent titanium, from about 0 to about 0.5 percent columbium, from about 0.25 to about 2.0 percent molybdenum, from about 0 to about 0.15 percent hafnium, and the balance nickel plus incidental impurities, the superalloy having a phasial stability number  $N_{V3B}$  less than about 2.10.

Advantageously, this superalloy composition may be further comprised of (percentages are in weight percent) from about 0 to about 0.04 percent carbon, from about 0 to about 0.01 percent boron, from about 0 to about 0.01 percent yttrium, from about 0 to about 0.01 percent cerium and from about 0 to about 0.01 percent lanthanum. Although incidental impurities should be kept to the least amount possible, the superalloy can also be comprised of from about 0 to about 0.04 percent manganese, from about 0 to about 0.05 percent silicon, from about 0 to about 0.01 percent zirconium, from about 0 to about 0.001 percent sulfur, and from about 0 to about 0.10 percent vanadium. In all cases, the base element is nickel. Furthermore, this superalloy can advantageously have a phasial stability number N<sub>V3B</sub> less than about 1.85, and a chromium content of from about 1.8 to about 3.0 percent, a rhenium content of from about 5.5 to about 6.5 percent, and a cobalt content of from about 2.0 to about 5.0 percent. This invention provides a superalloy having an increased resistance to creep under high stress, high temperature conditions, particularly up to about 1975°F.

Single crystal articles can be suitably made from the superalloy of this invention. The article can be a component for a turbine engine and, more particularly, the component can be a gas turbine blade or gas turbine vane.

The preferred superalloy compositions of this invention have a critically balanced alloy chemistry which results in a unique blend of desirable properties. These properties include: excellent single crystal component castability, particularly for moderately sized blade and vane components: adequate cast component solutionability: excellent resistance to single crystal cast component recrystallization; ultra-high creep-rupture strength to about 1975°F; extremely good low cycle fatigue strength: extremely good high cycle fatigue strength; high impact strength; very good bare hot corrosion resistance; very good bare oxidation resistance; and adequate microstructural stability, such as resistance to the undesirable, brittle phases called topologically close-packed (TCP) phases.

Accordingly, it is an object of the present to provide superalloy compositions and single crystal articles made therefrom having a unique blend of desirable properties. It is a further object of the present invention to provide superalloys and single crystal articles made therefrom for use in advanced gas turbine engines under high stress, high temperature conditions, such as up to about 1975°F These and other objects and advantages of the present invention will be apparent to those skilled in the art upon reference to the following description of the preferred embodiments.

#### BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a chart of hot corrosion test results performed to 117 hours on one embodiment of the alloy of this invention and on two prior art alloys;

FIG. 2 is a chart of hot corrosion test results performed to 144 hours on another embodiment of the alloy of this invention and on a prior art alloy.

#### DESCRIPTION OF THE PREFERRED EMBODIMENTS

The preferred nickel-based superalloy of the present invention comprises the following elements in percent

45	by weight:	
	Rhenium	about 5.0-7.0
	Chromium	about 1.8-4.0
	Cobalt	about 1.5-9.0
	Tantalum	about 7.0-10.0
50	Tungsten	about 3.5-7.5
	Aluminum	about 5.0-7.0
	Titanium	about 0.1-1.2
	Columbium	about 0-0.5
	Molybdenum	about 0.25-2.0
55	Hafnium	about 0-0.15
	Nickel + Incidental Impurities	balance

Nickel + Incidental Impurities balance

This superalloy composition also has a phasial stability number N<sub>V3B</sub> less than about 2.10. Further, this invention has a critically balanced alloy chemistry which results in a unique blend of desirable properties. These

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properties include increased creep-rupture strength relative to prior art single crystal superalloys, single crystal component castability, cast component solutionability, single crystal component resistance to recrystallization, fatigue strength, impact strength, bare hot corrosion resistance, bare oxidation resistance, and microstructural stability, including resistance to TCP phase formation under high stress, high temperature conditions.

Unlike prior nickel-based superalloys known in the art, the superalloys of the present invention have a low chromium, low cobalt and high rhenium content. The chromium is about 1.8-4.0% by weight. Advantageously, the chromium content is from 1.8% to 3.0% by weight. This chromium content is significantly lower than that typically found in prior art single crystal nickel-based superalloys. In the present superalloy, chromium provides hot corrosion resistance, although it may also assist with the alloy's oxidation capability. Tantalum and rhenium also assist toward hot corrosion property attainment, and aluminum is present at sufficient levels to provide adequate oxidation resistance, so that relatively low addition of chromium is tolerable in this alloy. Besides lowering the alloy's gamma prime solvus, chromium contributes to the formation of Cr, Re, W-rich TCP phase and must be balanced accordingly in these compositions.

The cobalt content is about 1.5-9.0% by weight. Advantageously, the cobalt content is from 2.0% to 5.0% by weight. This cobalt content is lower than that typically found in prior art single crystal nickel-based superalloys. In the present superalloy, cobalt assists in providing an appropriate heat treatment window since it has the effect of lowering the alloy's gamma prime solvus while generally not affecting its incipient melting point. Rhenium-containing alloys are generally designed with much higher cobalt content than the present invention for the purpose of imparting increased solid solubility and phasial stability. However, the superalloys of the present invention unexpectedly show that much lower cobalt contents are possible and desirable toward providing optimized phasial stability, including control of TCP phase formation.

The rhenium content is about 5.0-7.0% by weight and, advantageously, rhenium is present in an amount of from 5.5% to 6.5% by weight. The amount of rhenium in the superalloy of the present invention is significantly greater than the rhenium content of prior art single crystal nickel-based superalloys. Furthermore, the superalloys of this invention are generally designed with an increased level of refractory element content, e.g., W+Re+Mo+Ta. The tungsten content is about 3.5-7.5% by weight and, advantageously, the amount of tungsten is from 3.5% to 6.5% by weight. Tungsten is added since it is an effective solid solution strengthener and it contributes to strengthening the gamma prime. Additionally, tungsten is effective in raising the alloy's incipient melting temperature. The amount of tungsten added to these superalloys is balanced with the amount of rhenium added since they both contribute to the formation of "freckle" defects during the single crystal investment casting process. They also both strongly affect the propensity for TCP phase formation.

Similar to tungsten, rhenium is effective in raising the alloy's incipient melting point. However, rhenium is a more effective strengthener than tungsten, molybdenum and tantalum in terms of elevated temperature creep-rupture and, therefore, rhenium is added appropriately. Additionally, rhenium has a positive influence on this alloy's hot corrosion resistance. Moreover, rhenium partitions primarily to the gamma matrix, and it is effective in slowing gamma prime particle growth during high temperature, high stress conditions; Besides requiring the balancing of rhenium with tungsten for castability reasons, W+Re must also be set at a level consistent with minimizing TCP phase formation. In general, the TCP phases which occur in such material are rich in chromium, tungsten, and rhenium content, with rhenium being present in the greatest proportion. Thus, careful Re/W ratio control is necessary in this alloy to control the propensity for TCP phase formation.

The molybdenum content is about 0.25-2.0% by weight. Advantageously, molybdenum is present in an amount of from 0.25% to 1.5% by weight. Molybdenum is a good solid solution strengthener, but it is not as effective as tungsten, rhenium and tantalum. However, since the alloy's density is always a design consideration, and the molybdenum atom is lighter than the other solid solution strengtheners, the addition of molybdenum is a means of assisting control of the overall alloy density in the compositions of this invention.

The tantalum content is about 7.0-10.0% by weight and, advantageously, the tantalum content is from 8.0% to 10.0% by weight. Tantalum is a significant contributor to this alloy's strength through means of solid solution strengthening and enhancement of gamma prime particle strength (tantalum also partitions to the gamma prime phase). In this alloy, tantalum is able to be utilized at relatively high concentration since it does not contribute to TCP phase formation. Additionally, tantalum is an attractive single crystal alloy additive in this composition since it assists in preventing "freckle" defect formation during the single crystal casting process. Tantalum is also beneficial in this composition since it tends to raise this alloy's gamma prime solvus, and it is effective toward promoting good alloy oxidation and hot corrosion resistance, along with aluminide coating durability.

The aluminum content is about 5.0-7.0% by weight. Furthermore, the amount of aluminum present in this composition is advantageously from 5.3% to 6.5% by weight. Aluminum and titanium are the primary elements comprising the gamma prime phase. These elements are added in this composition in a proportion and ratio consistent with achieving adequate alloy castability, solution heat treatability, phasial stability and high mech-

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anical strength. Aluminum is also added to this alloy in proportions sufficient to provide oxidation resistance.

The titanium content is about 0.1-1.2% by weight. Advantageously, titanium is present in this composition in an amount from 0.2% to 0.8% by weight. Titanium is generally beneficial to the alloy's hot corrosion resistance, but it can have a negative effect to oxidation resistance, alloy castability and alloy response to solution heat treatment. Accordingly, the titanium content must be maintained within the stated range of this composition.

The columbium content is about 0-0.5% by weight and, advantageously, the columbium content is from 0 to 0.3% by weight. Columbium is a gamma prime forming element and it is an effective strengthener in the nickel-based superalloys of this invention. Generally, however, columbium is a detriment to alloy oxidation and hot corrosion properties, so its addition to the composition of this invention is minimized. Moreover, columbium is added to this invention's composition for the purpose of gettering carbon, which can be chemi-sorbed into component surfaces during non-optimized vacuum solution heat treatment procedures. Any carbon pick-up will tend to form columbium carbide instead of titanium or tantalum carbide, thereby preserving the greatest proportion of titanium and/or tantalum for gamma prime and/or solid solution strengthening in this alloy.

The hafnium content is about 0-0.15% by weight and, advantageously, hafnium is present in an amount from 0.02 to 0.05% by weight. Hafnium is added in a small proportion to the present composition in order to assist with coating adherence. Hafnium generally partitions to the gamma prime phase.

The balance of this invention's superalloy composition is comprised of nickel and small amounts of incidental impurities. Generally, these incidental impurities are entrained from the industrial process of production, and they should be kept to the least amount possible in the composition so that they do not affect the advantageous aspects of the superalloy. For example, these incidental impurities may include up to about 0.04% by weight manganese, up to about 0.05% by weight silicon, up to about 0.01% by weight zirconium, up to about 0.001% by weight sulfur, and up to about 0.10% by weight vanadium. Amounts of these impurities which exceed the stated amounts could have an adverse effect upon the resulting alloy's properties.

Additionally, the superalloy may optionally contain about 0-0.04% by weight carbon, about 0-0.01% by weight boron, about 0-0.01% by weight yttrium, about 0-0.01% by weight cerium and about 0-0.01% by weight lanthanum.

Not only does the superalloy of this invention have a composition within the above specified ranges, but it also has a phasial stability number  $N_{V3B}$  less than about 2.10. Advantageously, the phasial stability number  $N_{V3B}$  is less than 1.85 and, preferably, the phasial stability number  $N_{V3B}$  is less than 1.75. As can be appreciated by those skilled in the art,  $N_{V3B}$  is defined by the PWA N-35 method of nickel-based alloy electron vacancy TCP phase control factor calculation. This calculation is as follows:

#### **EQUATION 1**

Conversion for weight percent to atomic percent:

Atomic percent of element  $i = Pi = \frac{Wi/Ai}{\Sigma i (Wi/Ai)} X100$ 

where:

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Wi = weight percent of element i Ai = atomic weight of element i

#### **EQUATION 2**

Calculation for the amount of each element present in the continuous matrix phase:

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Element Cr R<sub>Cr</sub>=0.97P<sub>Cr</sub>-0.375P<sub>B</sub>-1.75P<sub>C</sub>

Ni 
$$R_{H_{1}}^{=P_{H_{1}}+0.525P_{B}-3(P_{Al}+0.03P_{Cr}+P_{I_{1}}-0.5P_{C}+0.5P_{V}+P_{Ia}+P_{Cb}+P_{H_{1}})}$$

Ti, Al, B, Ri=0

C, Ta, Cb, Hf

V Rv=0.5P<sub>V</sub>

W  $^{*}R_{(M)}=P_{M}-0.167P_{C}$   $P_{M}$ 
 $P_{M0}$   $P_{M}$ 

Mo  $R_{(M0)}=P_{(M0)}-0.75P_{B}-0.167P_{C}$   $P_{M0}$ 

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\*Note: weight percentage Re is added to weight percentage W for the calculation above.

#### **EQUATION 3**

Calculation of  $N_{V3B}$  using atomic factors from Equations 1 and 2 above:

$$N_i^i = \frac{R_i}{iR_i}$$
 then  $N_{v3B} = \Sigma_i Ni(N_v)i$ 

where:

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i = each individual element in turn.

N<sub>i</sub>I = the atomic factor of each element in matrix.

(N<sub>v</sub>)I = the electron vacancy No. of each respective element.

This calculation is exemplified in detail in a technical paper entitled "PHACOMP Revisited", by H. J. Murphy, C. T. Sims and A. M. Beltran, published in Volume 1 of International Symposium on Structural Stability in Superalloys (1968), the disclosure of which is incorporated by reference herein. As can be appreciated by those skilled in the art, the phasial stability number for the superalloys of this invention is critical and must be less than the stated maximum to provide a stable microstructure and capability for the desired properties under high temperature, high stress conditions. The phasial stability number can be determined empirically, once the practitioner skilled in the art is in possession of the present subject matter.

The preferred superalloy of this invention can be used to suitably make single crystal articles, such as components for turbine engines. Preferably, this superalloy is utilized to make a single crystal casting to be used under high stress, high temperature conditions characterized by an increased resistance to creep under such conditions, particularly high temperature conditions up to about 1079°C (1975°F). While this superalloy can be used for any purpose requiring high strength castings incorporating a single crystal, its particular use is in the casting of single crystal blades and vanes for gas turbine engines. This alloy possesses an unusual resistance to component recrystallization during solution heat treatment, which is considered an important alloy characteristic that is necessary when producing advanced technology, multi-piece, cast bonded single crystal airfoils. Additionally, this superalloy provides the alloy castability characteristics believed necessary to produce conventional-process-cast, moderately-sized turbine airfoils with intricate cooling passages.

While this superalloy's primary use is in aircraft turbine engines, there are stationary engine applications requiring the specialized high performance characteristics of this alloy. This is particularly the case in turbine engines which require performing characteristics with very restricted clearances, thereby materially limiting the amount of permissible creep. Engines designed to develop high performance characteristics are normally operated at higher component temperatures and, therefore, the problem of creep is increased. Generally, creep in excess of 1% is considered unacceptable in these cases. The creep characteristics of known state of the art alloys have limited operating temperatures and, thus, maximum performance capability. The superalloy of this invention has an increased resistance to creep under high stress, high temperature conditions, particularly up to 1079°C (1975°F).

The single crystal components made from this invention's compositions can be produced by any of the single crystal casting techniques known in the art. For example, single crystal directional solidification proc-

esses can be utilized, such as the seed crystal process and the choke process.

The single crystal castings made from the superalloy of the present invention are advantageously subjected to a high temperature aging heat treatment in order to optimize the creep-rupture properties of these alloys. This invention's single crystal castings can be aged at a temperature of from about 1066°C (1950°F) to about 1163°C (2125°F) for about 1 to about 20 hours. However, as can be appreciated by those skilled in the art, the optimum aging temperature and time for aging depends on the precise composition of the superalloy.

This invention provides superalloy compositions having a unique blend of desirable properties. These properties include: excellent single crystal component castability, particularly for moderately sized blade and vane components; excellent cast component solutionability; excellent resistance to single crystal cast component recrystallization ultra-high creep-rupture strength to about 1079°C (1975°F) extremely good low cycle fatigue strength; extremely good high cycle fatigue strength; high impact strength; very good bare hot corrosion resistance; very good bare oxidation resistance; and microstructural stability, such as resistance to formation of the undesirable TCP phases. As noted above, this superalloy has a precise composition with only small permissible variations in any one element if the unique blend of properties is to be maintained.

In order to more clearly illustrate this invention and provide a comparison with representative superalloys outside the claimed scope of the invention, the examples set forth below are presented. The following examples are included as being illustrations of the invention and its relation to other superalloys and articles, and should not be construed as limiting the scope thereof.

#### **EXAMPLES**

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A large number of superalloy test materials were prepared to investigate the compositional variations and ranges for the superalloys of the present invention. Some of the alloy compositions tested and reported below fall outside the claimed scope of the present invention, but are included for comparative purposes to assist in the understanding of the invention. Representative alloy aim chemistries of those materials tested are reported in Table 1 below.

#### TABLE 1

																A HYAV	⇔ Belo	
ALLOY		<u> </u>	В	Cr	<u>_c</u> o_	Мо	Ä	Cb Ti	۸١	Ţa	Re	Вf	Ni	NV38*		2_	3	
CMSX-10A				3.0	8.5	.70	7.2	.30 .65	6.0	7.6	5.0	.05	BAL	2.08		6.65	14.55	
-10B				2.6	8.2	.70	6.95	.30 .68	6.0	7.9	4.95	.06	BAL	2.02	11.9	6.68	14.88	
-10C		-	-	2.5	7.7	.70	6.6	.30 .65	5.9	8.2	4.8	.05	BAL	1.90	11.4	6.55	15.05	
-100			-	4.0	4.8	.60	6-4	.30 .60	5.7	8.2	4.9	.03	BAL	1.95	11.3	6.30	14.80	
-10E			-	2.2	7.2	.70	6.3	.25 .72	5.85	8.3	4.8	.042	BAL	1.84	11.1		15.12	
-10F	. 0	Ź	.02	2.4	7.6	.65	6.45	.28 .63	5.9	8.5	5.0	.046	BAL	1.89		6.53	15.31	
- 10G	٠.		-	2.4	6.3	.50	6.4	.20 .70	5.8	8.0	5.5	.04	BAL	1.82	11.9	6.5	14.7	
-10Ga	, .		-	2.4	4.0	.50	6.2	. 15 . 55	5.8	8.3	5.6	.04	BAL	1.72	11.8	6.35	14.8	
10K-10Gb			-	2.3	3.3	.40	5.5	.10 .30	5.7	8.4	6.3	.03	BAL	1.60	11.8	6.0	14.5	
-10H			-	2.2	5.9	.50	6.4	.15 .80	5.9	8.0	5.5	.04	BAL	1.82	11.9	6.7	14.85	
-101			-	2.5	4.7	.50	6.4	.15 .70	5.8	7.9	6.0	.04	BAL	1.81	12.4	6.5	14.65	20.9
-101			-	2.5	3.3	.40	6.1	.10 .60	5.8	7.9	6.0	.04	BAL	1.69	12.1.	6.4	14.4	20.4
· -10J			-01		4.0	.50	6.0	.20 .65	5.8	9.0	5.5	.04	BAL	1.79	11.5	6.45	15.65	
-10L	-	•	•	2.0	2.7	.40		.10 .20	5.65	8.4	6.3	.03	BAL	1.50	11.6	5.85	14.35	20.4
CMSX-12A	,			3.0	4.5	.35	5.5	- 1.0	5.65	9_0	5.5	.04	BAL	1.84	11.0	6.65	15.65	20.3
-128				3.5	3.0	.35	5.0	90	5.60			.04	BAL	1.80	11.0	6.5	15.3	20.19
-12C				2.8	3.5	.40	5.3	75	5.60			.04	BAL	1.70	11.1	6.35	15.15	20.3
120-12Ca			-	2.5	3.2	.45	4.7	50	5.60			.03	BAL	1.61	11.0	6.10	14.8	20.15
-12E			-	2.0	3.0	.45	4.7	40	5.60			.03	BAL	1.50	11.0	6.0	14.7	20.15
CMSX-10Ri			. <b>.</b>	2.65	7.0	.60	6.4	.40 .80	5.8	7.5	5.5	.06	BAL	1.91	11.9	6.6	14.5	20.0
CMSX-12Ri				3.4	8.0	.50	6.0	-				.06		1.92			14.6	19.4
											-						•	
	K	ey:	:		+ Re													
				_	l + .T	-												
. •				_			. + Cb											
				4 - W		+ Mo												

Third generation single crystal alloy development to investigate the compositional variations for the superalloys of the present invention began with the definition and evaluation of a series of experimental compositions. Increased creep-rupture strength was the primary objective of the initial development effort, with elemental balancing to provide a combination of useful engineering characteristics following the definition of a base concept for increased strength.

The initial materials explored the utility of higher levels of refractory element and gamma prime forming elements than are present in similar prior art compositions. As shown in Table 1, the alloy chromium content was reduced to improve alloy stability. Cobalt content, initially thought to be required for increased solid solubility, could be significantly reduced. Refractory element content (W+Re+Mo+Ta) was varied, while the summation of the primary gamma prime partitioning elements (Al+Ti+Ta+Cb) was also varied. The alloy's Re content was initially explored at conventional levels, but it was found that the Re level had to be increased.

Standard  $N_{\rm V3B}$  calculations were performed during the initial alloy design stage to assist respective alloy phasial stability predictions, with that number varying from one alloy composition to another.

Some of the alloys were produced using production-type procedures. These alloys were vacuum induction melted in the Cannon-Muskegon Corporation V-1 furnace, yielding approximately 91-136kg (200-300 lbs.) of bar product per alloy (see Table 2 below). Quantities of each compositional iteration, as reported in Table 2, were made into test bars and test blades by vacuum investment casting. Solution heat treatment procedures were developed in the laboratory in 76 and 152 mm (3" and 6") diameter tube furnaces. Gamma prime aging treatments were also performed in the laboratory.

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TABLE 2

Y-1 VIN FURRACE HEAT CHEMISTRIES

	Alley	Heat No.	C	В	Cr	Co	Ко	v	Cb	11	AL	To	Re	HE H
10 15	CNSX-10A -103 -1081 -1081 -1081 -1281 -1281 -1281 -106a -126 -100b (-10x) -120a (-120)	VF 778 VF 831 VF 965 VF 966 VF 963 VF 963 VF 964 VF 979 VF 983 VF 985 VF 985 VF 994 VF 994	.001 .002 .001 .001 .001 .001 .001 .001	<.001 <.001 <.001 <.001 <.001 <.001 <.001 <.001 <.001 <.001 <.001 <.001	2.9 2.6 2.65 2.69 2.66 3.3 3.4 3.4 2.4 2.7 2.2	8.5 8.2 7.0 7.0 7.0 8.0 8.0 8.0 3.95 3.5 3.3	.7 .6 .6 .6 .48 .50 .41 .45 .40	7.2 6.9 6.4 6.3 6.3 6.0 6.1 6.1 5.3 5.5 6.8	.3 .3 .4 .4 .4 <.05 <.05 <.05 .14 <.05 .09 <.01	.70 .68 .80 .79 1.01 1.00 1.00 .56 .75	6.05 6.06 5.72 5.66 5.78 5.69 5.60 5.36 5.83 5.66 5.74	7.6 7.6 7.6 7.6 7.6 7.6 8.4 8.8 8.2 8.6	5.0 5.5 5.4 5.3 5.3 5.3 6.4 6.4	.05 BA .05 BA .06 BA .06 BA .07 BA .06 BA .06 BA .03 BA .025BA .025BA

All other specimens reported in Table 1 above were produced by blending base alloy bar stock with the virgin elemental additions necessary to achieve the desired composition. The blending was done during test bar and blade manufacture. The base alloy bar stock plus virgin additions were placed into the casting furnace melt crucible, melted and the bath homogenized prior to pouring into an appropriate shell mold. It is believed that good correlation between alloy aim chemistry and test bar/blade chemistry was routinely achieved (see Table 3 below).

TABLE 3.

ALLOY TEST BAR CHEMISTRIES

Alloy	с	В	Cr	Co	Xo_	U	СР	Ţi	A.L	<u>Ta</u>	Яe	Hf	иt	¥~3B*
CTIST														2.09
CHSX-10A	-	-	2.9	8.5	.68	7.4	.29	.69	6.0	7.5	5.1	.07	BAL	2.01
- 108	-	-	2.7	8.1	.69	6-95	. 29	.69	6.0	7.8	4.8	.06	BAL	1.86
-10C	٠.	-	2.6	7.7	.69	6.4	.30	.62	5.7	8.3	4.7	.07	BAL	1.83
- 100	-	-	4.0	5.0	.62	6.0	.31	.59	5.44	8.1	4.7	.04	BAL	1.84
- 10E	-	-	2.2 .	7.2	.70	6.4	.26	.63	5.89	8.2	4.8	.05	BAL	1.86
- 10F	.014	.027	2.4	7.7	.65	6.4	.28	.63	5.96	7.9	5.0	.04	BAL	
- 106	•	-	2.5	6.5	.53	5.5	. 20	.68	5.6	8.2	4.6	.05	BAL	1.68
- 10Ga	-	-	2.4	4.0	.41	6.2	. 14	.55	5.79	8.3	6.0	.025	BAL	1.73
-10Gb (10K)		-	2.3	3.5	.42	5.9	.10	.43	5.67	8.5	6.0	.024	BAL	1.63
- 10H		-	2.3	5.6	.51	6.2	.17	.76	5.58	7.8	5.4	. 05	BAL	1.69
- 101	-	-	2.6	4.8	.52	6.6	.14	.67	5.65	7.4	5.4	.04	BAL	1.70
• 10 La	_	-	2.7	3.5	.47	5.2	. 10	.60	5.80	8.0	5.8	.04	BAL	1.67
- 101 a	.017	01	2.6	4.0	.48	6.0	. 19	.62	5.74	8.8	5.7	-04	BAL	1.76
- 101		-												
- 12A		_	3.0	4.6	.39	5.3	<.01	.96	5.61	9.4	5.0	.05	BAL	1.80
- 128		_	3.5	3.0	.38	5.1	<.01	.84	5.52	8.8	6.1	.05	BAL	1.79
		-	2.7	3.5	,45	5.4	<.01	.75	5.62	8.8	6.0	-04	BAL	1.72
- 12C			2.5	3.2	.46	5.0	<.01	.61	5.56	8.7	6.0	.03	BAL	1.60
-12Ca (120)	•	-	2.,	J.L		J	- • • •							
- 12E	•	-	2 45	7.0	.60	6.4	.40	.80	5.67	7.6	5.5	.065	BAL	1.87
- 10R I	-	•	2.65		.48	6.1	<.01	.99	5.54	7.6	5.3	.07	BAL	1.92
- 12R i	-	-	3.4	8.0	.40	0.1	1.01	.,,	3.54	• • • •		-		

<sup>.</sup> PWA H-35 Hethod

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For the CMSX-10D specimen (see Table 1), high quality virgin elemental additions were vacuum melted

and the refined material was poured into 51 mm (2") diameter bars. In turn, a quantity of the resulting bar was used to produce single crystal test bar/blade specimens by investment casting.

It was apparent that considerable variation in the investment casting process integrity may have occurred during specimen manufacture since varying levels of test bar freckle formation, secondary dendrite arm spacing and property attainment were apparent. Derivative alloy response to solution treatment (reported in Table 4 below) varied, and was a function of both alloy composition and test specimen quality.

Heat treatments developed for the alloy iterations are reported in Table 4 below. Full gamma prime solutioning was desired for each material, however, this objective was not universally achieved. Primary gamma prime aging was performed to effect a more desirable gamma prime particle size and distribution. Secondary gamma prime aging was performed to effect precipitation of conventional matrix gamma prime precipitates along with ultra-fine gamma prime precipitates located within the matrix channels between the primary gamma prime particles for these specimens.

#### TABLE 4

20				Heat Treat	ment Detail	
20	Atloy	Peak Solution	on Temperature	x & r Solutioned*	Primary Y' Aging+	Secondary d' Aginge
		• 5	<u>.c</u>			
	444	2460	1349	97.0-98.0	1975°F/4 Hrs	1600°F/20 + 1400°F/24
25	CMSX-10A -108	2465	1352	97.0-98.0	1975°F/4 Hrs 1975°F/19.5 Hrs	1600°F/20 + 1400°F/24
	- 1DC	2470	1354	99.0-99.5	2100°F/8 Hrs	1600°F/20 + 1400°F/24
	*		47/7	99.9-100	1975°F/10 Hrs 1975°F/10 Hrs	1600°F/22 + 1400°F/24
	-100	2450 2465	1343 1352	100	1975*F/15 Hrs	1600°F/20 + 1400°F/24
	-10E	2403	1372		1975°F/21 HF#	1600°F/25.5 + 1400°F/23
	-10F	2444	1340	95	1975°F/16 Hrs	1600°F/23 + 1400°F/24 1600°F/24.5 + 1400°F/17
30	-106	2475	1357	99.0-99.5	1975°F/12 Hrs	1600°F/20 + 1400°F/23
	-10Ga	Z485-90	1363-65	99.5-100	2075°F/5 Hrs 2075°F/6 Hrs	1600°F/24 + 1400°F/30
					20/3 //6 11/8	1612°F/48 + 1414°F/22
	4001 44040	3/05	1363	100	2075°F/6 Hrs	1600°F/24 + 1400°F/30
	-10Gb(10K) -10H	2485 2475	1357	98.5-99.0	1975°F/16 Hrs	1600°F/27.5 + 1400°F/27
	- jun	. 2773			1975"F/18 Hrs	1600°F/101 + 1400°F/46
	- 101	2475	1357	100	2075°F/5 Hrs	1600°F/22 + 1405°F/24 1600°F/24 + 1400°F/24
35	-101 s	2480	1360	99.5-100	2075°F/5 Hrs 1975°F/15 Hrs	1600°F/24 + 1400°F/30
	- 10J	2480	1360	98.0-99.0	2075°F/5 Hrs	1600°F/24 + 1400°F/30
	-10L			98.5-99.0	1975°F/16.5 Hrs	1600°F/24 + 1400°F/32
	<b>-12</b> A	2475	1357	98.3-17.0	1975°F/12 Krs	1600°F/24 + 1400°F/27.5
		2480	1360	99.0-99.5	1975°F/13 Hrs	1600°F/57 + 1400°F/39
	-12B -12C	2485-90	1363-65	99.5-100	2075*F/5 Hrs	1600°F/20 + 1400°F/5
40	-120	2403-70	1305 03		2075°F/6 Hrs	1600°F/24 + 1400°F/30
	-12Ca(12D)	2485	1363	100	2075"F/6 Hrs	1600°F/24 + 1400°F/30
	-12E			98.5-99.8	2075°F/6 Hrs	1600°F/24 + 1400°F/30
	-10Ri	2460	1349	98.5-99.8 100	2075°F/6 Hrs	1600°F/24 + 1400°F/30
	-12R i	2455	1346	100	<b>2017 17 9</b>	

\* Determined by visual estimation

+ Specimens air cooled from all aging treatments

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Fully heat treated test bars were creep-rupture tested. The specimens were machined and low-stress ground to ASTM standard proportional specimen dimension. The specimens were creep-rupture tested at various condition of temperature and stress, according to standard ASTM procedure.

A significant factor of the CMSX-1OA alloy design was the shift to higher Re content. At the sane time, W, Cr, Ta and other gamma prime strengtheners were balanced to provide the desired alloy characteristics and properties. The alloys higher Re level resulted in significantly improved creep-rupture strength throughout the entire test regime, as indicated by the results reported in Table 5 below for the CMSX-10A specimens.

TABLE 5

DISX-104 CREEP-PUPTURE

				PAINT TAILS				M HOURS
10			_	¥	FINAL CRE	EP READING		REACH
		MENTURE TIME	% ELDNG.	. RA	t, hours	% deformation	1,0%	2.0%
	TEST COMOITION	HOURS	ELLMG.			-		21.0
			24.2	26.9	534.2	22.331	10.9	8.7
	1600*f/75.0 ks1	534.4	22.0	27.8	328.3	21.055	6.3	72.2
	871°C	328.4	21.1	26.3	526.3	17.552	28.4	12.4
	071 0	527.3	21.1	20.5	•			
			•	34.5	304.2	28.614	62.1	108.9
15	1700°//50.0 ksi	305.0	31.1	19.9	291.8	19.324	71.5	123.7
	927°C	292.4	19.2		85.7	1-474	65.9	•
	321 C	87.6	2.6	5.8	67.7	•••		
					413.8	15.643	182.7	246.1
	1800°F/30.0 ksi	415.6	16.1	21.4		34.326	460.4	524.3
		848.0	37.1	33.0	846.3	32.984	476.8	655.1
	982°C	1016.Z	33.2	30.5	1014.3	32.904	4.0.0	
							395.0	425.0
20		586.5	38.1	38.0	585.6	33.050	395.0	422.0
20	1800°F/36_0 ksi	572.7	36.9	35.3	570.7	29.029	373.0	406.0
	982°C:	546.5	26.4	34.2	545.7	25.843		317.6
			22.4	26.3	418.7	18.105	286.7	353.2
		420.3	14.8	17.0	425.1	10.244	326.5	
		426.0	24.3	23.8	239.7	23.264	94.1	123.9
		239.8		27.4	253.6	18.510	115.2	152.7
		255.7	19.9	27.7	2,50.0			
	•			11.0	31.0	2.075	26.7	30.7
25	1900°F/25.0 ksl	32.3	5.5		128.7	39.556	30.4	48.1
	1038°C	129.7	43.2	38.9	166.1	30.816	58.2	78.4
	1030 €	168.7	34.7	36.4	226.4	16.926	146.3	160.6
		228.1	18.1	32.3		27.323	9.9	29.9
		277.7	29.5	31-1	276.4	35,121	218.4	250.9
		423.4	39.7	38.3	422.7	34.861	192.9	226.7
		383.8	35.9	36.1	382.7		211.6	238.0
		373.3	31.3	35.7	371.6	26.138	.,,,,	•
30	•	313.3					33.9	77.0
		138.0	22.3	33.0	136.3	19.052	54.7	71.9
	2000°F/18.0 ksi	134.9	40.7	36.5	134.7	38.328	50.1	69.4
	1093°C	122.9	23.2	34.9	122.0	19.050	40.B	56.8
			34.2	36.6	114.4	30.861		157.9
		115.6	35.1	36.2	244.3	29.844	135.7	140.0
		245.2	36.3	35.4	221.8	33.737	113.0	61.4
	*	221.9		34.2	180 - 1	29.249	53.1	01.4
35		181.2	32.1	J-1.E				69.8
33				49.0	124.1	30.086	45.8	
	2050°F/15.0 ksi	126.4	47.9	47.8	148.1	39.308	16.8	34.5
	1121°C	150.5	45.5	40.0	138.7	23.596	30.6	76.4
	112)	140.5	30.6		120.0	29.479	16.3	55.6
		120.8	29.5	39.7	79.0	11.644	41.7	54.8
		79.0	11.7	14.4	112.1	21.401	55.9	69.5
		112.2	24.3	31.3	112.1	21.401		
						20.520	42.2	62.6
40	747095/47 E bai	94.1	22.1	27.5	94.1	20.320	28.0	58.8
	2100°F/12.5 ksi 1149°C	112.5	39.4	33.1	112.2		52.3	62.5
	1145 C	96.6	25.9	35.9	95.5	14.542	40.9	63.5
	•	123.6	43.4	40.4	122.9 .	31.050		37.6
			21.7	29.6	49.9	9.330	35.1	. 38.5
		50.8	41.6	43.7	89.7	37.422	13.6	- 30.7
		90.5	41.0	73.1				286.0
				35.1	419.9	23.196	213.8	
45	1800°F/36.0 ks1 *	420.6	23.9	34.0	394.7	31.623	239.4	264.9
		396.1	37.1	34.0	382.9	25.554	220.5	247.9
	982°C	384.9	31.1	34.0	300.7			
				*				

As-Solutioned Condition

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Microstructural review of the failed rupture specimens of this alloy revealed that TCP phase precipitation occurred during the respective creep-rupture tests, particularly those at  $1038^{\circ}$ C ( $1900^{\circ}$ F) and above. It became apparent that the  $N_{v38}$  phasial stability number calculation would be an effective tool in predicting alloy

stability and, effectively, high temperature creep strength for the invention.

Wherein the CMSX-10A specimen's  $N_{v3B}$  number was 2.08, CMSX-10B was designed to the 2.02 level. This was accomplished by the further reduction of alloy Cr content and similar reduction to Co and W+Re level. W was reduced more than the Re in this specimen since Re is more effective in the solid solution. Additionally, wherein some loss in W contribution to the gamma prime could be anticipated, it was sufficiently replaced by the modest increase to Ta content in this composition. These changes resulted in the CMSX-10B alloy specimen exhibiting even more improved creep strength at 982°C (1800°F) Table 6 reported below illustrates that three specimens achieved an average life of 961 hours, with 1.0% creep occurring at an average of 724 hours. However, it was observed that TCP phase was present at higher temperature.

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#### TABLE 6

#### CHESK-108 CREEP-RUPTURE

1800°F/40 982°C 1900°F/40 1900°F/25 1038°		907.1 989.3 988.4 507.0 598.1 408.3 265.3 385.3 412.8 389.3 459.5	19.2 18.9 35.9 44.1 46.9 62.6 39.7 45.5 43.4 51.5	34.0 33.5 36.1 45.4 43.4 52.1 43.7 46.2	907.0 988.5 987.3 505.7 596.1 407.2 262.7	17.832 17.657 31.813 41.548 42.340 54.479	697.2 768.1 705.8 317.9 386.5 187.3	752.7 817.8 767.5 352.6 415.2
1800° f /40 982°( 1900° f /25 1038'		989.3 988.4 507.0 598.1 408.3 265.3 385.3 412.8 389.3 459.5	18.9 35.9 44.1 46.9 62.6 39.7 45.5 43.4	33.5 36.1 45.4 43.4 52.1 43.7	988.5 987.3 505.7 596.1 407.2	17.657 31.813 41.548 42.340 54.479	768.1 705.8 317.9 386.5	817.8 767.5 352.6 415.2
1800° f /40 982°( 1900° f /25 1038'		989.3 988.4 507.0 598.1 408.3 265.3 385.3 412.8 389.3 459.5	18.9 35.9 44.1 46.9 62.6 39.7 45.5 43.4	33.5 36.1 45.4 43.4 52.1 43.7	988.5 987.3 505.7 596.1 407.2	17.657 31.813 41.548 42.340 54.479	768.1 705.8 317.9 386.5	817.8 767.5 352.6 415.2
1800° f /40 982°( 1900° f /25 1038'		989.3 988.4 507.0 598.1 408.3 265.3 385.3 412.8 389.3 459.5	18.9 35.9 44.1 46.9 62.6 39.7 45.5 43.4	33.5 36.1 45.4 43.4 52.1 43.7	987.3 505.7 596.1 407.2	31.813 41.548 42.340 54.479	705.8 317.9 386.5	767.5 352.6 415.2
1800° f/40 982°( 1900° f/25 1038'	C	968.4 507.0 598.1 408.3 265.3 385.3 412.8 389.3 459.5	35.9 44.1 46.9 62.6 39.7 45.5 43.4	36.1 45.4 43.4 52.1 43.7	987.3 505.7 596.1 407.2	41.548 42.340 54.479	317.9 386.5	352.6 415.2
982°0 1900°1/25 1038°		507.0 578.1 408.3 265.3 385.3 412.8 389.3 459.5	44.1 46.9 62.6 39.7 45.5 43.4	45.4 43.4 52.1 43.7	505.7 596.1 407.2	41.548 42.340 54.479	386.5	415.2
982°0 1900°1/25 1038°		598.1 408.3 265.3 385.3 412.8 389.3 459.5	46.9 62.6 39.7 45.5 43.4	43.4 52.1 43.7	596.1 407.2	42.340 54.479	386.5	
982°0 1900°1/25 1038°		408.3 265.3 385.3 412.8 389.3 459.5	62.6 39.7 45.5 43.4	52.1 43.7	407.2	54.479		
982°0 1900°5/25 1038°		265.3 385.3 412.8 389.3 459.5	39.7 45.5 43.4	43.7				256.5
982°0 1900°5/25 1038°		385.3 412.8 389.3 459.5	45.5 43.4			37.102	87.6	119.2
982°0 1900°5/25 1038°		412.8 389.3 459.5	43.4		383.5	39.031	177.4	213.4
982°0 1900°1/25 1038°		389.3 459.5		40.5	410:6	38.771	189.1	233.4
982°0 1900°1/25 1038°		459.5		44.2	386.8	36.920	220.5	249.2
982°0 1900°1/25 1038°			40.0	46.3	458.0	39.513	210.2	291.1
982°0 1900°1/25 1038°		258.9	38.1	40.6	257.9	36.743	32.	90.2
982°0 1900°1/25 1038°		484.1	27.9	40.0	483.4	26.296	288.1	326.7
982°0 1900°1/25 1038°		376.9	16.4	20.4	376.8	16.088	96.0	226.6
982°0 1900°1/25 1038°		481.0	50.5	48.2	478.8	34.557	264.4	297.5
982°0 1900°1/25 1038°		461.5	35.1	40.6	460.1	30.786	181.1	265.3
982°0 1900°1/25 1038°		483.0	47.1	46.8	482.1	43.714	286.2	320.7
982°0 1900°1/25 1038°					499.7	30.486	11.9	280.1
982°0 1900°1/25 1038°		500.1	33.4	37.0	477.7	30.400	,	
982°0 1900°1/25 1038°		436.7	40.2	44.1	436.2	39.818	294.6	318.9
1900*1/25		390.8	50.1	42.8	390.3	41.817	250.9	276.2
1038	С	336.9	52.7	48.1	335.2	46.697	226.5	240.9
1038		330.7	32.1	40.1	237.2	40.07		
1038	i n kai	237.8	55.9	45.7	237.4	53.854	33.0	113.5
		295.7	57.4	49.1	295.6	46.592	123.7	170.9
	i°C	£//	27.4	****	,			
	B.O ksi	192.7	31.5	26.6	191.6	27.733	56.3	88.6
1093		166.5	41.4	25.3	166.5	34.102	46.2	72.7
1093	, .	173.3	36.6	27.0	171.4	31.481	24.0	66.1
•								
2050°F/15	S.O ksi	219.6	40.1	40.4	218.6	37.871	13.2	56.8
2000 1,10		122.3	28.2	47.9	120.6	26.614	37.0	63.7
1121	0.0	118.4	33.2	60.0	116.9	29.986	36.7	56.5
1 (2)	C	179.7	44.1	48.1	179.1	39.188	8.4	75.3
	•	74.9	44.2	48.6	74.6	34 . 800	6.8	14.5
		168.3	48.6	49.7	. 167.0	43.171	36.9	77.1
		104.8	17.0	27.2	102.8	1.626	66.1	-
		155.9	46.3	49.8	155.2	38.388	64.4	81.9
		90.6	15.1	21.4	87.1	1.046	75.5	-
	•	120.5	46.3	55.8	118.7	35.143	10.3	27.7
		150.7	39.8	49.7	150.1	33.903	21.4	60.9
		149.5	33.2	46.2	148.9	23.166	73.3	88.3
		142.9	42.0	47.5	142.5	41.524	54.9	70.5
								•
2050°F/15	i.O ksi	163.0	52.5	49.2	161.9	46.146	20.5	76.9
		151.1	66.4	45.6	150.7	59.115	52.7	75.5
1121	, C	131.8	57.3	44.4	131.5	48.310	26.3	57.1
,		* 156.0	54.4	41.0	155.9	45.502	55.5	78.3
		* 133.7	57.2	56.0	132.7	41.753	67.5	80.7
		• 135.1	59.7	52.3	134.3	46.317	54.9	71.5
		151.1	66.4	45.6	150.7	59.115	52.7	75.5
		131.8	57.3	44.4	131.5	48.310	26.3	57.1
		131.0						
2100*1/15				48.1	10.1	43 434	25 7	36.3
1149	: O bef	69.7	54.Z	90. l	69.4	47.674	25.3	20.3

\* As-Solutioned Condition

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*5*0

Only about 97-98% gamma prime solutioning was achieved in the CMSX-10A and -10B materials (see Table 4) which was insufficient for the purpose of optimizing alloy mechanical properties and microstructural homogeneity. Attainment of a greater level of gamma prime solutioning, therefore, became an equal priority in

tandem with improving microstructural stability at temperatures above 1038°C (1900°F).

To confirm the suspected composition of the TCP-phase forming in the alloys, scanning electron microscope (SEM) wavelength dispersive x-ray (WDX) microchemistry analyses of CMSX-10B test bar contained needles was undertaken and compared to the alloys gamma and gamma prime compositions. The results, reported in Table 7 below, confirm that the needles were enriched in Cr, W and Re.

#### TABLE 7

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#### CMSX-108 Micro-Chemistry Analyses

- Cast Test Bar (VF 831)
- · Transverse Section. Bottom Bar Location.
- Solutioned to \$465°F) 1352°C
- Aged 1975°F/19.5 Hrs./AC 1079°C 1600°F/20 Hrs./AC - 871°C 1400°F/24 Hrs./AC - 760°C

				1400	F/24 Hrs./	AC -	. 760 C			
		GAHMA	PHASE		GAMMA PR	HE PHA	SE		NEEDLE CO	NSTITUENT
20	ELEN	K Z	A F	ELEM	K	2 A	F	ELEH	K Z	A F
	ALK	0.0101 1.090	0.324 1.000	ALK	0.0145 1.	.084 0.3	22 1.000	ALK	0.0116 1.107	0.347 1.000
	TIK	0.0069 1.007	0.930 1.051	TIK	0.0084 1.	.002 0.9	34 1.052	TIK	0.0077 1.026	0.908 1.039
	CRK	0.0428 1.008	0.963 1.108	CRK	0.0250 1.	.002 0.9	65 1.117	CRK	0.0390 1.028	0.949 1.083
	COK	0.0970 0.994	0.984 1.018	COK	0.0761 0.	988 0.9	87 1.022	COK	0.0755 1.016	0.977 1.025
	NIK	0.6891 1.033	0.988 1.010	NIK	0.7270 1.	026 0.9	91 1.005	MIK	0.6143 1.056	0.983 1.024
	TAL	0.0485 0.794	1.020 1.000	TAL	0.0697 0.	788 1.0	24 1.000	TAL	0.0389 0.814	1.018 1.000
25	W L	0.0329 0.788	0.963 1.000	WL	0.0311 0.	783 0.9	62 1.000	W L	0.0682 0.808	0.968 1.000
	REL	0.0422 0.785	0.968 1.000	REL	0.0085 0.	779 0.9	68 1.000	REL	0.1083 0.805	0.973 1.000
			WT X				uit x			VT X
	ELEM	CPS	ELEM	ELEH	CPS		ELEH	ELEH	CPS	ELEM
30	AL K	12.1800	2.87	AL K	17.940	ю	4.19	AL K		3.02
	TI K	5.5200	0.71	TI K	6.840	10	0.86	TI K		0.79
	CR K	27.6400	3.98	CR K	16.450	Ю	2.31	CR K	21.5800	3.69
	CO K	40.6800	9.74	CO K	32.540	10	7.64	CO K	27.1700	7.42
	NI K	253.1300	66.84	RI K	272.380	0	71.11	NI K	193.7500	57.84
	TA L	6.5667	5.99	TA L	9.632	9	8.64	TA L	4.5259	4.70
	¥ L	4.0775	4.33	W L	3.937	5	4.13	WL	7.2620	8.71
	RE L	4.6000	5.56	RE L	0.950	0	1.13	RE L	10.1300	13.82
35										
	TOTAL	•	100.00	TOTAL	<u>.</u>		100.00	TOTAL	•	100.00

The calculated N<sub>v38</sub> numbers were 1.90 for CMSX-10C and 1.95 for CMSX-10D. Re was maintained at around 5% while W was further reduced to improve stability in these specimens. Alloy Ta was increased since it did not participate in TCP formation and the Ta/W ratio was effectively improved, which assisted with alloy castability. Chromium was reduced in the -10C specimens but increased to 4.0% in the -10D specimens to provide an opportunity to determine the suitability of the Cr levels from a hot corrosion standpoint. Co was reduced in both materials, significantly in the -10D specimen, while Al+Ti level was also reduced to assist in achieving more complete gamma prime solutioning. Creep-rupture results for the two specimens are reported below in Tables 8 and 9, respectively. Even though the -10D alloy specimens were observed to exhibit full gamma prime solutioning (as opposed to 99.-99.5% for CMSX-10C) the alloys greater Cr content, which necessitated a lower Al+Ti level, effected lower properties than attained with CMSX-10C. However, both materials exhibited improved alloy stability and higher temperature properties, so that attempts to balance the alloys low and high temperature creep response were favorable.

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#### TABLE 8

## DISX-10C CREEP-RUPTURE

								REACH
	•	PLIPTURE TIM	E X	¥		EEP READING	1.0%	2.0%
10	TEST CONDITION	ROURS	ELONG.	RA	t, hours	X deformation	1.04	
	1,41							
		*** 4	31.4	30.5	555.2	26.615	316.1	376.3
	1800°F/36.0 ksi	556.1		37.5	636.4	38.460	416.6	455.4
	982°C	636.6	43.9	34.7	607.6	19.074	410.6	460.6
	•	609.2	23.3			34.991	407.3	443.4
		635.7	44.9	45.6	635.3		409.8	438.7
		612.8	43.5	38.8	611.9	41.951	407.0	-55
15							61.1	166.3
	1850°F/36.0 ksi	252.2	30.2	37.8	252.0	22.033		194.8
	•	298.1	41.3	39.0	297.6	37.953	170.3	146.0
	1010°C	231.1	33.6	39.5	230.2	29.689	127.8	140.0
		492.4	52.5	52.4	491.6	48.922	176.5	251.7
	1922°F/20.3 kei		38.6	45.5	528.9	33.353	269.6	306.2
	1050°C	529.8	48.9	43.3	635.2	45.804	189.5	318.3
20	1030 €	637.5	40.7	. 43.3				
20				41.5	258.7	32.444	74.2	127.5
	2000°F/18.0 ksi	258.8	35.0		292.1	42.079	145.6	170.9
	1093°C	293.1	49.2	44.1	220.9	33.507	55.6	123.3
	1093 C	221.9	43.0	48.5		33.759	113.6	143.6
	•	266.1	35.1	44.0	264.6	33.777	113.0	
						27.755	26.0	134.8
	2050°F/15.0 kel	196.6	39.7	40.3	194.1		11.1	51.4
	<del>-</del>	170.4	30.1	46.3	169.2	25.624	46.5	76.5
25	1121°C	193.2	38.1	42.9	191.9	32.288		150.8
		247.3	33.1	40.5	246.0	26.494	122.0	1,50.0

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#### rable 9

## DISK-100 CREEP-RUPTURE

40	TEST CONDITION	MUPTURE TIME	I ELONG.	X RA	FINAL CRI t, hours	EP READING I deformation		REACH Z.OX
	1800°F/36.0 ksi	425.0	26.7	29.3	426.3	24.166	189.2	24B.3
45	982°C 1850°7/36.0 ksi 1010°C	141.0 140.7 166.0	23.1 14.7 17.5	26.8 26.1 28.9	140.1 140.2 165.0	20.660 13.741 15.640	57.8 56.2 76.5	77.7 77.6 100.1
45	1922°F/20.3 kei 1050°C	519.9 667.0 680.3	23.8 17.6 14.9	24.9 23.7 28.2	518.9 665.2 678.9	22.608 16.819 14.476	202.0 151.8 340.2	345.6 391.4 500.3
50	2000°F/18.0 ksi 1193°C	370.3 - 401.5 366.6	18.8 11.1 17.5	21.3 18.0 25.8	369.9 400.0 366.6	15,560 8,903 8,049	20.9 — 19.8 223.9	106.9 125.5 306.1
••	2050°F/15.0 ksi 1121°C	465.3 338.8	12.9 9.8	20.5 24.8	465.2 337.7	12.639 9.468	61.0 30.8	305.9 204.4

The acceptability of the alloys' low Cr content was confirmed through extremely aggressive short-term burner rig hot corrosion tests performed at 899°C (1650°F) 1% sulfur, 10 ppm sea salt condition. FIGS. 1 and 2 illustrate the results for tests performed to 117 and 144 hours for the CMSX-10C and CMSX-10D specimens, respectively. In both cases, the materials performed similar to MAR M 247-type materials, thereby confirming the suitability of the low Cr alloy design concept.

With the above-noted results, another series of alloys, CMSX-10E, -10F, -10G, -10H, -10I, and -12A were designed, produced and evaluated. The alloys explored Re level ranging 4.8-6.3%, 2.2-3.0% Cr level, 4.7-7.6% Co level and the remainder balanced to maintain castability, improve solutionability and improve phasial stability. The N<sub>v3B</sub> number ranged between 1.81-1.89.

One of the series, CMSX-10F, contained .02% C and .02% B. These additions were observed to improve casting yield and may have assisted in providing more consistent control of single crystal cast article orientation. However, the melting point depressants, C and B, restricted the specimen's response to solution heat treatment. The CMSX-10F creep-rupture properties are reported in Table 10 below.

TABLE 10

2	Y	1	ì	
•	۰	۰	•	

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				CKSX-1UF	CKEEP-KOP TOKE	i		
	TEST CONDITION	RUPTURE TIME HOURS	X ELONG,	Z RA	FINAL CRE t, hours	EP READING % deformation		IN HOURS O REACH 2.0%
25	1800°F/36.0 kel 982°C	616.0 666.6 603.1	18.1 45.6 25.3	22.4 48.0 24.3	615.8 666.4 602.5	16.898 43.261 24.281	439.9 464.6 398.4	477.6 492.3 444.0
	1850°F/36.0 kef 1010°C	243.9 285.9 258.6	19.6 26.8 19.2	28.2 32.1 29.1	243.0 285.5 258.3	18.045 25.701 18.175	129.1 187.8 168.3	160.9 206.0 189.5
30	1922°F/20.3 kel 1050°C	499.5 649.2 361.0	40.0 55.6 15.8	41.0 52.9 21.9	498.5 648.3 357.7	37.756 51.045 2.599	208.2 197.6 273.2	272.6 338.8 335.7
	2000°F/18.0 ksi 1093°C	235.4 276.1 290.0	39.6 43.7 36.7	51.7 52.8 47.3	235.4 274.4 289.1	37.881 36.762 33.304	100.8 115.1 125.3	133.2 155.9 162.1
35	2050°f/15.0 kei 1121°C	255.4 255.1 254.5	28.7 33.4 25.4	36.6 43.1 33.3	255.0 254.9 254.4	27.426 31.378 23.737	67.4 46.2 50.9	131.0 102.2 118.7

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The CMSX-10E, G, H and I, plus CMSX-12 creep-rupture specimen results are reported below in Tables 11, 12, 13, 14, and 15, respectively. The results show a general improvement to alloy creep-rupture strength aboven 1038°C (1900°F) while maintaining extremely good strength at lower temperatures.

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## TABLE 11

## CHSX-10E CREEP-RUPTURE

		-						IN HOURS
		RUPTURE TIM	E X -	x	FINAL CRE	EP READING	1	O REACH
	TEST CONDITION	HOURS	ELONG.	RA	t, hours	1 deformetion	1.0%	2.0%
10								
10								
	100015 /36 0 hei	664.5	31.4	36.3	663.5	30.435	436.5	470.8
	1800°5/36.0 ks i 982°C	604.4	35.1	36.7	603.3	33.371	· 253.7	355 <i>.9</i>
	702 C	582.5	41.5	36.1	581.7	39.792	78.9	329.3
		553.5	35.9	37.0	552.5	33.172	326.4	357.1
		257.9	25.3	32.0	257.0	22.734	149.4	170.3
15	1850°F/36.0 ksi			32.1	198.6	16.261	122.4	139.4
15	1010°C	. 199.2	18.4		259.7	31.315	159.9	174.0
		260.5	33.6	33.4	234.1	31.373	137.7	
	1922°F/20.3 ksi	810.6	38.6	33.0	508.4	33.523	210.2	378.2
	1050°C	800.9	35.3	36.4	799.1	32.405	339.7	434.2
	1050 C	859.9	39.0	35.4	859.6	37.036	364.6	465.2
		2/2 8	27.7	29.3	362.4	24.887	98.4	177.3
	2000°F/18.0 ksi	362.8		27.0	409.9	26.426	173.6	218.6
20	1093°C	411.2	29.4		368.8	12.941	170.3	221.9
		369.7	15.3	28.2	379.2	27.656	177.9	206.6
		379.7	26.4	26.1	3/7.2	27.050	*****	20010
	:2050°F/15.0 kmi	476.9	21.8	23.4	. 476.3	18.233	196.6	255.9
		418.4	27.5	24.7	417.5	25.854	180.0	227.3
	1121°C	397.7	19.0	23.8	396.8	17.522	112.6	198.2
	1	*****						

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## TABLE 12

## DISK-10G DREEP-RUPTURE

10		MUPTURE TIM	E 1	x	FINAL CRE	EP READING		IN NOURS O REACH
	TEST COMDITION	HOURS	ELONG.	ŘA.	t, hours	X deformation	1.0%	2.0%
	TEST CONTITON		<u> CLUMO.</u>					
							447.2	508.1
	1700°F/55.0 ksi	671_8	19.6	28.6	670.5	14.775	441.2	493.4
	927°C	693.6	26.0	24.2	691.7	21.750 19.913	464.8	520.4
		724.9	23.3	29.7	723.2	15.200	77.0	356.7
15		582.5	18.6	20.1	581.1	19.115	56.4	314.8
		681.2	20.9	24.1	679.2	17.857	242.1	308.7
		538.4	21.6	17.5	538.3	14.157	235.3	308.0
		523.0	17.7	21.8	522.4 568.5	15.035	287.0	354.9
		569.7	17.5	19.8	306.3	15.035	201.0	234
	**************************************	775.2	29.6	29.3	773.8	28.826	315.0	539.9
	1800°F/36.0 ksi	719.7	29.5	28.5	717.8	27.266	321.2	486.4
20	982°C	741.6-	28.0	25.9	740.3	24.870	284.5	464.2
20		682.8	45.6	34.7	681.1	39.289	409.1	452.4
		764.0	23.2	33.7	764.0	22.884	543.6	586.6
		790.4	41.4	35.6	789.4	38.172	511.6	565.3
		799.1	27.0	32.3	797.4	25.737	529.8	579.1
		.,,	27.00					
	1850°F/36.0 ksi	354.4	19.3	30.2	351.9	16.000	246.7	271-4
	1010°C	344.5	28.5	31.9	344.3	26.174	220.8	241.9
25	10.0 €	315.4	23.7	30.7	315.1	23.571	183.4	205.6
								462.1
	1922°F/20.3 ks1	753.4	31.7	34.8	753.2	27.914	352.3 281.1	402.1
*	1050°C	728.0	31.5	33.5	727.1	28.362 30.770	339.3	437.3
		731.6	34.3	38.8	730.5	30.770	337.3	431.3
				29.4	94.9	22.842	41.5	50.9
	1976°F/28.1 kei	. 95 . 4	29.3	27.2	94.7	20.130	45.8	54.7
30	1080°C	95.7 104.6	26.7 30.4	33.2	104.4	27.517	41.8	54.4
30		100.8	25.6	35.1	98.9	21.577	49.2	58.1
		95.8	25.9	28.9	93.6	19.748	41.1	51.4
		110.0	29.3	30.3	108.0	22.669	48.5	60.1
		108.2	27.3				43.8	
		104.8					45.8	
		104.3					48.6	
						•		
35	2000°F/18.0 ksi	464.4	23.1	21.3	463.6	18.190	257.7	293.5
	· ·	411.9	18.3	23.0	410.4	16.347	103.5	227.6
	1093°C	370.9	27.0	38.7	369.8	25.326	7.6	47.3
								404.0
	2012"F/14.5 ksi	790.2	31.2	34.9	788.7	24.939	299.9	406.0
	1100°C	671.4	23.6	25.7	670.3	13.397	303.3	396.3 277.7
	1100 C	512.1	22.6	28.1	510.4	21.094	192.5	434.7
40		651.7	27.4	39.7	651.3	16.328	315.7	388.7
40		754.6	29.7	25.4	753.1	24.032	193.8	300.7
		908.3	17.7	18.3	-:-		388.7	438.2
		758.9	30.8	26.5	758.7	24.090	300.7 316.5	426.7
		740.0	19.8	20.5	739.5	16.962	316.5 359.8	412.4
		671.5	26.4	23.8	669.3	15.578	226.5	272.2
	ZO50°F/15.0 ksi	410.8	22.9	27.4	410.0	18.655	156.4	191.2
	1121°C	283.5	18.0	31.2	283.5	15.303 12.979	156.4	191.2
45	1121 C	320.0	16.8	17.4	318.3	18.488	29.9	189.1
		283.5 320.0 389.7 381.4	. 22.0	22.1	389.7 381.1	24.758	69.5	197.9
	4	381.4	27.0	24.1	301.1	24.730	07.13	*****
		251 1	12.7	30.4	252.9	8.984	108.4	185.5
	2100°F/12.0 ksi	254.4 419.8	20.5	26.0	419.8	18.917	201.1	274.3
•	1149°C	4 17.0	20.5	20.0				
50	2100°F/12.5 ks1	331.4	16.9	21.7	331.1	15.069	25.2	83.2
<b>5</b> U	1149°C	367.7	19.2	23.2	366.5	17.530	76.2	177.4
	1147 C	387.3	16.8	17.2	386.5	12.742	236.9	282.0
		383.1	34.1	32.4	381.6	32.135	10.5	164.3

## TABLE 13

## DISK-10H CREEP-RUPTURE

								IN HOURS
	TEST COMOTITION	BLPTURE TIME	¥	X		FINAL CREEP READING		REACH
10		HOURS	ELONG.	RA_	t hours	X deformation	1.0%	2.0%
	_		77 7	27.2	563.2	Z2.669	318.5	366.2
	1800"f/36.0 ks l	563.4	23.2 24.5	23.0	552.7	21.324	373.1	402.8
	982°C	553.1		27.3	526.4	19.715	358.2	390.7
		526.9	20.7	41.4	594.4	32.090	328.8	372.8
		594.5	35.1	•1.•	377.7	55111		
15		7/7 0	24.3	20.1	242.2	20.686	107.3	155.6
	1850"F/36.0 ks i	242.9	17.0	21.0	221.0	14.888	115.9	150.4
	1010°C	221.9 223.4	21.3	21.0	221.7	19.196	128.4	144.7
		223.4	21.3					
		520.6	26.1	29.3	520.4	23.183	234.3	319.1
	1922°F/20.3 ks1		26.3	21.2	469.2	19.333	176.1	253.2
	1050°C	470.4 574.7	16.8	23.0	573.0	14.411	282.1	373.0
		3/4./	10.0					
20		434.0	21.5	18.7	432.1	20.234	103.5	233.1
	2000°F/18.0 ks1	437.3	27.1	33.8	437.3	126.306	182.6	240.8
	1093°C	430.7	24.6	20.4	430.7	23.244	68.8	192.1
		430.1	21.1	19.3	428.9	19.D50	73.7	213.B
		430.1						
		366.1	16.3	12.0	365.5	11.326	239.8	273.3
	2050°F/15.0 kei	384.0	17.4	16.0	382.3	12.055	168.2	242.9
	1121°C	420.2	12.2	13.3	418.6	10.017	127.3	273.2
25								

TABLE 14

## CHESK-10 | CREEP RUPTURE

35		RUPTURE TIME	1	x	FUNAL CRE	EP READING	TIRE IN	
-	TEST COMOTTION	HOURS	ELONG.	RA"	t, hours	% deformation	1.0%	2.0%
	1800°F/36.0 ksl	565.1	35.2	32.0	564.8	29.774	297.0	368.9
	982°C	581.9	32.4	29.3	580.2	28.689	371.9	402.5
		514.1	24.1	30.2	514.1	21.207	318.3	358.2
	1850/36.0	260.5	25.0	24.8	259.3	23.255	156.7	175.3
40	1010°C	247.5	22.4	29.1	245.7	17.730	131.9	169.0
40	1010 €	246.1	23.7	29.0	246.1	20.277	137.6	156.7
	1922/20.3	916.3	24.9	30.3	914.8	22.465	472.9	549.3
	1050°C	934.8	32.2	33.0	934.8	30.165	353.7	475.2
	.050 0	863.6	27.8	28.5	862.9.	27.057	295.6	442.5
	1976/28.1	116.1	19.5	20.1	116.1	19.155	57.4_	70.1
	1080°C	65.6	22.9	20.6	64.2	21.368	17.8	26.4
45		91.6	23.2	25.3	90.4	15.544	37.6	49.7
	2000/18.0	430.1	22.7	25.7	429.2	18.449	58.9	193.0
	1093°C	483.8	19.8	25.1	483.8	17.860	102.4	245.4
	2050/15.0	397.7	17.9	30.0	. 397.3	13.264	239.8	292.9
	1121°C	487.7	21.4	21.9	487.1	18.854	248.2	318.4
	1121 C	468.3	18.4	25.5	467.9	15.800	194.1	300.1
50	2100/12,0	501.3	10.1	15.9	498.7	0.615		-
	1149°C	401.3	16.8	26.3	399.7	15.429	6.6	25.5
	1149 C	210.6	11.5	12.7	210.3	0.373	•	-

#### TABLE 15

#### CHSX-12A CREEP-RUPTURE

		SUPTURE TIME X				FINAL CREEP READING		TIME IN HOURS TO REACH	
	TEST CONDITION	HOLKS	ELONG.	X RA	t hours	% deformation	1.0%	2.0%	
10	1800°F/36.0 ks1	491.9	40.2	41.6	491.8	38.605	254.0	293.7	
10	982°C	420.4	23.5	31.9	420.3	19.299	234.9	277.9	
		383.4	25.3	26.2	382.9	22.920	198.1	244.3	
		456.2	24.1	26.1	454.5	22.582	89.9	265.5	
	•	458.0	30.7	32.7	457.1	26.155	253.2	292.8	
	•	386.8	30.1	30.4	386.3	27.031	172.7	216.9	
		403.7	34.5	28.8	402.7	31.033	140.2	204.9	
		398.7	21.6	23.5	398.4	20.277	181.1	236.1	
15	1850/36.0	208.5	32.1	40.5	208.3	31.248	100.8	119.6	
	1010°C	189.5	21.2	25.2	189.4	20.461	99.1	116.3	
	1922/20.3	829.6	46.5	45.3	828.8	44.488	315.8	400.7	
	1050°C	797.0	33.5	32.5	796.9	32.856	315.3	400.5	
	2000/18.0	500.3	31.7	29.6	499.2	24.922	218-4	268.5	
	1093°C	227.6	36.5	41.2	227.1	26.825	90.6	113.9	
20	.055 @	430.4	18.5	23.3	430.4	18.180	181.0	234.1	
	2050/15.0	424.8	17.0	27.5	423.3	15.832	263.5	301.2	
	1121°C	366.1	26.2	42.8	365.5	20.399	146.6	197.8	
	1121 6	400.8	18.2	25.4	400.7	16.910	184.6	251.3	
	2100/12.0	255.4	25.8	45.8	253.6	22.920	64.1	125.B	
	1149°C	483.9	10.1	19.3	482.7	8.602	378.6	421.9	
25	1143 C	325.1	7.1	16.6	324.7	4.315	268.8	302.5	

Varying the primary gamma prime aging treatment was explored with most of the development activity concentrated on achieving optimized gamma prime size and distribution through longer soak times at 1080°C (1976°F) see Table 4) since higher temperature aging treatments accelerated TCP phase formation during the aging cycle.

Ten to twenty-one hour soak times at 1975°F were successful since they provided average gamma prime particles of about 0.5 um dimension. However, it appeared that shorter primary gamma prime aging time at higher temperature may be more practical, once more stable microstructures were defined.

Microchemical SEM WDX needle particle analyses was performed on a failed CMSX-10G creep-rupture specimen. The specimen, tested at 1976°F/28.1 ksi condition, exhibited needles in its microstructure. The results of the analysis are reported in Table 16 below and indicate, again, that the needles formed in this class of material are particularly rich in Re, but are also enrichened with Cr and W.

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#### TABLE 16

#### CMSX-10G

1080°C (1976°F)28.1 ksi 104.6 HRS.

ELEM	K	Z	A	F
CRK	0.0426	1.105	0.793	_
COK	0.0584	1.094	0.888	1.086
NIK	0.1740	1.140	0.910	1.116
WL	0.2107	0.941	0.972	1.000
REL	0.4767	0.941	0.979	1.000

NEEDL	E CHEMISIK	<u> </u>
•	•	WT %
ELEM	CPS	ELEM
CRK	113.7000	4.63
COK	112.1100	5.54
NIK	305.1425	15.02
WL	134.8988	23.03
REL	276.4000	51.76

A standardized test for resistance to recrystallization was performed on a CMSX-10G test bar. The test method and the results are reported in Table 17 below. The test results indicate that the CMSX-10G specimen exhibited similar resistance to cast process/solution treatment/bonding process recrystallization level in comparison to CMSX-4 alloy.

100.00

#### TABLE 17

## TABLE 17

Method: A controlled level of compressive stress is imparted on the entire surface of an as-cast test bar. The bar is then solution heat treated. Following solution treatment, the bar is sectioned and the transverse section is observed metallographically. Depth of recrystallization measurements are taken.

. Evaluation Standards:

Alloy	Depth of RX	Resistance To RX Anticipated in Blade Castings
CHSX-4 SX 792 CHSX-10G	.1016mm(.004") Entire Bar .1016mm(.004")	Very Good Very Poor Very Good

The CMSX-10Ga -10Ia, -12B,-12C, -10J, -10Ri and -12Ri compositions were defined and evaluated. No creep-rupture properties were generated for the CMSX-10J specimen, although test bars were produced and a solution heat treatment developed. Again, the inclusion of C and B in the -10J composition appeared to have

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positive effect to single crystal test specimen yield. Additionally, the lower levels of C and B than evaluated in CMSX-10F specimen, particularly lower B, made the material more amenable to solution heat treatment. Nine-level typical of the CMSX-10F composition.

The CMSX-10Ga and -10Ia alloys were designed with N<sub>v3B</sub> numbers of about 1.70. These alloy specimens contain about 2.5% Cr, 3.3-4.0% Co, 5.6-6.0% Re, greater Ta/W ratio, reduced Cb, and reduced Al+Ti content. Such reduction to Cb+Al+Ti level improved the solutioning characteristics of the materials (see Table 4), plus assisted achievement of increased alloy stability. Both specimens exhibited nearly full gamma prime solutioning.

The lowered N<sub>v3B</sub> number continued to show effectiveness in providing better creep-rupture capability at temperature greater tha 1038°C (1900°F), while maintaining extremely good creep-strength at lower temperature. CMSX-10Ga test results from specimens produced with improved casting process controls exhibited 700 hours or more life with about 475 hours required to creep to 1.0% for 982°C (1800°F) 36.0 ksi condition. For higher temperature exposure, the specimen provided the improved average life of about 500 hours at 1121°C (2050°F)/15.0 ksi condition and average 1.0% creep deformation that occurred at about 250 hours, as indicated by the results reported in Table 18 below.

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## TABLE 18

#### DICK-10GA CREEP-RUPTURE

	•	*					TIME IN	MOURS
	1621 CONDITION	MUPTURE TIME	II ELONG.	X RA	FINAL CREE t, hours	Y deformation	1.0%	2.0%
10			19.9	25.2	499.7	19.541	316.5	360.1
	1800°f/36.0 ksi.	500.7	19.9 29.1	25.4	583.9	26.395	370.0	401.8
	982°C	584.2 505.1	22.6	29.8	503.7	18.212	307.4	347.3
		730.9	42.0	42.8	730.7	40.216	477.6	516.1
		730.4	42.0	42.0			460.6 428.5	
15	1850/36.0	184.5	41.0	33.9	163.2	37.154 19.323	82.3 191.6	94.5 207.8
,,,	1010°C	291.5	27.3	29.9	290.2 278.1	29.054	155.3	180.5
	•	279.5	33.9 30.9	32.5 36.6	322.9	29.218	194.1	217.1
		323.9 326.5+	30.V 8.9	12.6	366.7		•	•
		295.2+	33.3	33.5	-	•	•	-
		277.2-	33.3				174.1 162.3	
		300.1+	22.8	22.4	•	•	102.3	-
20					4	~	39.7	48.9
	1976/28.1	88.6	34.9 28.2	33.9.	88.6	25.502 19.706	53.9	61.3
	1080°C	100.1	∴28.2	29.2 31.4	98.8 107.0	23.657	51.1	62.1
	1000 C	107.9	28.8 27.0	33.8	87.1	24.177	39.2	48.5
		87. 1 82. 8	23.3	27.7	81.0	17.301	20.6	38.0
		82.0 88.2	31.2	35.2	86.4	24.463	33.6	44.4
		83.7	34.0	34.3	83.4	29.718	₹36.3	45.1
25		114.1	24.3	26.3	113.0	20.544	62.1	73.2
		122.3	18.3	21.3	120.7	15.740	76.5	86.0
		117.7	23.2	25.6	117.5	22.284	78.0	B5.3
		. (INTERRUPTE	D TESTS)		40.2	1.036 1.187	39.9 42.3	-
			60.1	38.8	43.4	-	***	-
		99.3+	60. I 41.5	34.5	127.5	37.493	51.2	62.6
		127.9 96.8	22.9	27.9	96.5	20.126	45.9	54.4
30		118 0	31.3	27.1	118.0	24.603	49.5	61.3
		118.9	25.0	22.8	110.2	21.521	46.4	58.0
	•	96.6+	24.1	22.9	•	•	-	-
		120.5+	25.8	29.4	•	•	. •	•
		113.0+	27.6	20.5	•	· i.	•	•
	1976/18.85	(1NTERRUPTE	D 1E515)		261.5		260.3 204.6	-
35	1080°C				207.2	1.017	210.1	305.9
33	1000	592.1	25.8	22.4	590.4	23.596 26.289	293.3	332.6
		570.7	27.2	26.9	570.7 535.2	17.513	308.2	344.2
		535.5	19.3	23.9	337.2	17.713	240.5	
							307.6	
	****	536.8	28.5	27.5	535.6	20.662	232.3	321.3
	2050/15.0 1121°C	497.0	23.7	23.9	496.2	17.600	260.3	317.9
40	1121 C	514.8	23.4	24.4	513.1	12.500	230.4	340.4
		454.1	16.6	35.2	453.7	15.476	263.2	317.1
		420.3+	33.7	33.2	-	•	•	-
		(INTERRUP)		-	-	•	<b>Z39.</b> 1	-
				-	•	-	189.6	• •
		1.3		•	-	• .	280.3	
		560.1+	•	22.9	-	•	-	•
45	2012/14.5	536.6+	7.3	8.1				•
	1100°C	330,07	1.3	•••			424.6	
				7/ -	353.8	12.646	91.2	219.1
	2100/12.0	354.1	14.8	36.5	333.8	12.040	-	•
	1149°C	343.4+	•	27.2	•	=	91.4	
		ž.			•		147.2	
50	•					3		
					5	:		-
		491 0+	•	16.7	-	-		

<sup>1700/50.0 + 927°</sup>C + Machined From Blade Specimen

1% creep strength is a significant property. Limiting creep strains to 1.0% and 2.0% is extremely important to gas turbine component design, since a component's usefulness is generally measured by its resistance to creep to an approximate 1-2% level, not its ultimate rupture strength. Many prior art alloys may exhibit attractive rupture strength at the >1038°C (>1900°F) level, however, they lack the level of <u>useful</u> strength, i.e., creep strength to 2.0%, that this invention provides in tandem with its far superior strength in test conditions below 1038°C (1900°F).

The CMSX-10la specimens also provided significantly increased creep strength at the higher temperature extremes, but it did not appear to develop strength as good as the CMSX-10Ga specimens in lower temperature tests, as indicated by the results in Table 19 below.

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#### TABLE 19

#### CHSX-10 In CREEP RUPTURE

		RUPTURE TIME	x	· <b>x</b>	FINAL CREEP READING	TIME IN TO R	
	TEST CONDITION	HOURS	ELONG.	RA	t. hours % deformation	1.0%	2.0%
20	1800°F/36.0 kai 982°C	532.0 474.6 374.3	34.8 23.8 20.0	32.7 29.2 21.0	530.7 33.000 473.1 22.886 372.8 19.238	259.1 201.0 171.1	312.5 269.2 214.7
	1850/36.0 1010°C	256.0 251.4 217.8	28.7 34.4 30.5	28.5 30.3 22.4	256.0 27.867 250.7 33.055 217.2 27.000	135.4 121.6 94.2	157.1 144.6 117.9
25	1976/28.1 1080°C	85.7 81.9 68.9	27.5 33.6 26.1	28.9 31.8 25.8	83.8 21.754 81.0 24.384 67.6 20.960	36.9 32.1 23.1	46.2 42.1 32.4
•	2012/14.5 1100°C	930.2 844.4 864.2	10.0 17.7 15.3	14.4 23.2 11.9	928.4 9.649 842.8 16.132 862.8 14.558	104.6 339.7 179.9	455.7 502.3 453.4
30	2050/15.0 1121°C	510.2 528.6 438.8	17.8 17.9 14.3	19.7 24.2 11.3	508.4 15.703 527.0 14.873 436.4 13.556	187.2 293.7 56.0	312.7 364.3 136.9
	2100/12.0 1149°C	616.4 467.7	19.0 19.1	19.1 26.1	616.3 14.112 466.0 11.373	60.0 273.6	422.5 374.8

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Similarly, CMSX-12B, with Nv3B at 1.80 level and additional chemistry balance as presented in Table 1, provided attractive creep strength at test condition greater than 1038°C (1900°F), but did not perform quite as well as CMSX-10Ga in lower temperature tests, as indicated by the results reported in Table 20 below.

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TABLE 20

#### CHSX-128 CREEP-RUPTURE

		_		_		BEADING	TIME (N TO RI	
	TEST CONDITION	RUPTURË TIME HOURS	X ELOKG.	X RA	FINAL CREEP t hours	X deformation	1.0x	Z.0X
	1621 (000)1100					14.070	43.9	56.2
10	1976°F/28.1 ksi	91.7	15.3	17.2	91.2	17.396	27.4	36.8
	1080°C	72.6	19.4	23.2	72.6		8.6	11.9
	1080 C	14.1	5.0	1.3	12.7	2.300	17.8	38.9
		98.1	16.9	17.6	96.4	13.670	43.8	58.7
		108.2	25.2	24.1	108.0	22.794	46.1	60.1
		106.9	24.7	24.2	106.3	21.024	45.8	58.7
		104.8	24.0	26.8	104.3	20.094		60.8
		104.3	26.8	21.4	103.2	22.347	48.6	00.0
15		,						358.0
• -		515.0	24.7	24.2	513.3	19.468	320.1	
	1800/36.0		23.2	21.1	530.8	22.184	318.3	359.5
	982°C	536.4	13.2	19.9	302.9	12.582	166.0	200.8
	, JOE 0	304.7	13.2		, •		•	
			18.4	23.1	262.4	17.660	12.5	142.2
	1850/36.0	262.6	10.4	٠				
	1010°C		47.3	18.5	1029.5	15.113	428.0	703.7
20	2019/19°C	1031.3	17.2	20.0	1076.7	15.217	704.2	819.2
20		1078.7	15.6		839.2	9.282	607.6	677.7
	1100°C	839.4	14.9	22.8	834.8	18.024	591.1	658.5
		836.9	23.2	21.0	721.9	15.913	170.8	333.6
		722.0	16.4	21.1	710.8	12.490	381.9	531.5
		711.3	14.5	18.8		16.201	447.7	530.7
		711.9	18.3	20.0	711.4	10.201.	44771	
						9.394	70.4	360.4
	.2050/15.0	507.5	10.0	10.1	507.2		241.7	309.0
25		434.0	17.5	16.8	434.0	13.847	241.7	307.10
	1121°C	-5 -50					18.2	224.7
		487.5	25.3	20.3	486.6	20.986		413.6
	2100/12.0	444.9	7.8	11.0	442.2	3.884	347.3	713.0
	1149°C	777.7						

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Alloy composition has the greatest effect on ultimate creep strength. However, some of the variation experienced between alloy derivatives, and particularly for tests exhibiting inconsistent results for a given alloy, can be caused by variation in casting process condition. Casting process thermal gradient variation affects the cast specimen dendrite arm spacing and ultimately, its response to solution heat treatment and primary gamma prime aging treatment. It must, therefore, be recognized that much of the creep-rupture results reported herein may have been generated under non-optimized conditions and may be capable of improvement. Improved casting process control may provide casting microstructures more amenable to solution treatment and study to determine the appropriate primary gamma prime aging treatment to provide the optimum gamma prime particle size, which may result in further mechanical property enhancement.

The CMSX-12C composition was designed to provide a calculated N<sub>v3B</sub> number of 1.70. The alloy Cr content was designed at 2.8% and Co set at 3.5% aim for this alloy. An attractive Ta/W ratio was maintained while Re content was moderate at 5.8%. The alloy's Al+Ti content was reduced, in comparison to the CMSX-12A and CMSX-12B specimens, to provide improved alloy response to solution procedure.

Similar to the CMSX-10Ga specimen, the CMSX-12C specimen exhibited an improved balance of creep strength for test condition ranging 982-1149°C (1800-2100°F), as reported in Table 21 below.

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## CHSX-12C CREEP-RIPTURE

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	TEST CONDITION	RUPTURE TIME HOURS	ELONG.	X RA		P READING X deformation		IN HOURS REACH 2.0%
	1800-F/36.0 ksi	465.2	31.8	21.0	464.5	30.543		
	982°C	518.0	26.1	31.2	517.9	24.947	173.0 288.1	262.4 334.3
10		480.9	28.3	33.6	480.0	27.715	239.7	297.5
		713.3	30.0	28.0	713.2	28.899	455.0	503.7
	1850/36.0	237.7	28.2	26.8	237.7	27.054	114.4	145.3
	1010°C	221.2	22.9	27.3	220.7	22.491	111.3	135.2
		231.7 338.9	23.3 26.2	24.7	231.0	22.614	121.0	144.7
		300.1•	20.2	27.0	337.5	23.256	216.0	236.3
15		295.2•	33.3	33.5	•		÷	•
	1976/28.1	73.2	20.8	29.1	72.2	17.768	29.3	70.0
	1080°C	79.0	28.1	31.8	77.4	21.533	31.4	38.9 41.4
	.000 C	83.8	21.6	26.5	62.3	17.860	34.2	43.8
		67.6	31.2	29.8	67.5	24.177	25.5	34.6
	•	. 113.0+ 79.4						
20		76.2				•	30.8	
		68.8					32.8 29.3	
		118.1	26.0	28.0	116.2	23.822	49.3	62.0
						•	•	
		41			-	•	29.0	-
		(INTERRUPTED TEST	(2)		•	-	29.4	
25					•	-	32.9	-
	1976/18.85 ksi					•	65.4	
	1080°C						218.0	
	.000	*********			-	-	271.9	-
		(INTERRUPTED TESTS	•		•	•	168.9	-
	·				-	•	116.4 240.5	
30	2012/4/ 5						24013	
	2012/14.5	1001.8	23.6	20.0	1000.7	23.348	249.6	542.8
	1100°C	865.5	20.7	26.1	864.B	18.807	418.2	569.3
	1100 C	•	•				61.9 267.1	
	2050/15.0	509.4						
	1121°C	546.4	13.7 15.6	22.3 23.6	508.0	12.860	158.1	315.1
35		,,,,,,	13.6	23.6	546.4	14.044	323.0	404.0
			•				180.8 44.2	
						•	240.7	
							190.9	
	2100/12.0	404.3	11.2	21.6	404.3	8.438	290.1	326.4
	1149°C	321.7	9.5	15.0	320.4	7.671	156.6	254.1
40		545.1	8.2	22.1	542.2	5.351	236.0	452.9
		457.4	8.6	23.4	455.8	6.612	309.3	380.9
	2100+F/12.0	371.4+	14.2	17, 1	, <b>-</b>	•	-	-
	1750-149°C	*** 0.						
	954°C	446.9+	16.8	20.4	-	-	-	•
	1976•F/18.85	476.6+	19.2	27.1	_	_	•	
45	1080°C	459.94	30.6	30.2	<b>-</b>	•		
	1074-1129 1 1-2			•		• .		
	1976•f/28.1 ksi	120.5+	24.1	22.9	· •	•	-	-
	. 1080°C	99.64	25.8	29.4	-	-	•	•
	2050-f/15.0 ksi	469.8	-	30.8	_	_	_	
50	1121°C	485.4		22.7	-	•	-	•
<i></i>	2012•F/14.5 ksi	•						
							638.1 521.8	
	1100°C						267.1	
							61.9	
	.*						395.7	

Hachined from Blade Specimens

With improved casting process controls, this specimen has shown the following 1.0% longitudinal creep strengths, as reported in Table 22 below.

#### TABLE 22

Test Condition	Time to 1.0% Strain Hrs
982°C (1800°F)/36.0 ksi	455
1149°C (2100°F)/12.0 ksi	309.3

Both alloys provide similarly greater rupture strength than CMSX-4 alloy at condition to 1080°C (1976°F). Respective improvements to metal temperature capability are reported below in Table 23.

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#### TABLE 23

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	<u> Temperature</u>	Approx. Strength Advantage Relative to CMSX-4
20	982°C (1800°F)	22.2°C(40°F)
		25.0°C(45°F)
	1010°C(1850°F)	23.9°C(43°F)
	1080°C(1975°F)	25.5 0(.5 -7
25	Based on 1.0% creep stren	gth, the respective approximate
	982°C(1800°F)	+25.6°C(+46°F)
	1010°C(1850°F)	+33.3°C(+60°F)
	1010 C(1830 F) 1080°C(1976°F)	+30.6°C(+55°F)

Note that the comparison is not density corrected.

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For test temperature above 1080°C (1976°F), the test results indicate that the CMSX-10Ga and CMSX-12C specimens provided slightly lower strength than CMSX-4 alloy. The reduction in strength advantage for these alloys is believed to be the result of TCP phase formation. To address this issue, the alloys CMSX-10Gb, CMSX-10L, CMSX-12Ca, and CMSX-12E, are designed with N<sub>v38</sub> number as low as 1.50 (see Table 1) to provide greater phasial stability, and effect much improved high temperature creep-strength while maintaining most of the creep advantage demonstrated for the 982-1080°C (1800-1976°F) test regime.

The CMSX-10Ri and CMSX-12Ri compositions were designed at the 1.91 and 1.92 N<sub>V3B</sub> levels, respectively. These specimens were subjected to the most extensive testing of properties. They were designed with 2.65% and 3.4% respective Cr levels, with other features remaining similar to the aforementioned alloy design considerations. The properties generated for these two materials confirm the overall invention design concept with the other material iterations able to provide similar physical properties and relatively better blends of mechanical properties.

The CMSX-10Ri and CMSX-12Ri specimens' respective creep-rupture capabilities are reported below in Tables 24 and 25.

TABLE 24

## CMSX-10(Ri) CREEP-RUPTURE

		i	x	x	FINAL CREE	P PFADING	TIME IN TO RE	
10		MUPTURE TIME	ELONG.	ŘA	t. hours	X deformation	1.0%	2.0%
	TEST COMDITION	HOURS	ELUMU.				•	_
		227.3	21.2	33.8	225.4	14.359	52.8	131.5
	1675-1/75.0 ksi	231.6	19.3	31.0	231.3	16.671	51.0	125.1
	913°C	223.4	17.0	22.3	223.3	15.360	68.5	126.6
		263.4						
		425.9	18.3	33.7	425.6	16.047	303.4	334.7
15	1750/50.0 954°C	428.0	18.4	29.7	427.3	16.229	309.2	343.0 360.3
	954 C	460.8	17.1	25.7	459.0	15.308	314.7	364.3
							492.8	521.5
	1800/36 0	698.5	39.9	34.3	696.8	36.980	472.0	513.8
	1600/3&-0 982°C	676.3	28.3	33.3	674.5	27.221	469.3	504.9
	*	692.9	38.5	31.3	692.2	36.494	407.3	504.7
		•			291.1	31.774	194.1	210.4
20	1850/36.0	291.2	34.1	33.1		25.321	170.2	186.4
	1010°C	260.0	29.3	32.1	258.8	30.940	169.3	187.1
	.0.0	272.3	34.5	31.8	271.1	30.940	107.5	
					-613.5	50.482	365.8	415.5
	1850/27.56 1010 C	614.0	52.0	42.0 39.0	575.9	49.183	345.1	368.2
	1010°C	576.3	49.7	39.U 35.4	480.7	38.294	309.3	335.4
		481.1	40.4	33.4	400.7	30.274		
25	•		23.5	31.7	75.9	22.130	38.6	46.7
20	1976/28.1	76.2 80.5	19.0	26.3	79.8	14.665	44.3	51.3
	.1080°C	80.5 99.7	26.2	28.1	98.9	23.480	40.4	54.0
		· 99.7	20.2	20			41.4	-
		(INTERRUPT	EO 155151				37.0	-
		(INIERMO)	25 125.07		•		40.5	-
								184.8
30	1976/18.85	265.6	29.5	35.7	264.7	29.010	158.7	155.0
30	· · · · · · · · · · · · · · · · · · ·	278.8	51.4	38.8	278.1	46.026	82.0	
	1080°C	2.4.2	- : ·				139.7	-
		(INTERRUPT	ED TESTS)				128.8	-
		•••••					100.1	•
						77 /70	286.5	335.3
	2012/14.5	490.8	40.2	33.5	490.5	37.678 32.814	291.4	319.9
		447.0	37.0	41.5	445.0	32.814	113.5	• • • • • • • • • • • • • • • • • • • •
35	1100°C				-	• .	205.7	-
		(INTERRUP	ED TESTS)		-	•	202.2	•
					-	•	202.2	
					250.0	25.559	100.0	149.5
	2050/15.Q	251.9	33.6	35.9		23,149	177.5	221.2
	<sup>2050/15</sup> .8 1121°C	318.9	27.1	30.0	318.2	۵.147 -	181.0	•
					-		95.5	-
40					-		34.5	. •
		(INTERRU	PTED TESTS)	•	•			
		7	17.9	27.2	400-1	17.877	102.8	225.0
	2100/12.0	400.3	17.9	22.9	361.8	14.986	125.7	217.2
	1149°C	362.1	19.9	24.0	388.2	19.510	41.1	180.7
		389.5	17.7	27.0	200.00			
45								

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TABLE 25

#### CHEX-12(RI) CREEP-RUPTINE

							FIRE II	
		RUPTURE TIME	X	x	FINAL CREE			EACH
	TEST CONDITION	HOURS	ELDNG.	RA	t hours	% deformation	1.0%	2.01
		200 0	22.3	23.1	209.3	19.958	2.6	46.3
10	1675+F/75.0 ksi	209.8		17.4	189.7	12.483	1.6	42.5
	913°C	191.4	14.3		188.3	19.080	1.5	22.3
	715 C	189.6	22.0	22.8	188.3	14.000	1.5	22.3
	1750/50.0	448.1	26.7	26.6	447.9	26.054	302.3	335.5
		403.1	19.0	26.9	401.9	18.566	210.0	290.2
	954°C		19.4	26.9	434.4	18.503	89.1	284.1
		435.0	17.5	20.7	454.4	10.505		•
15	1800/36.0	604.5	34.7	29.9	604.3	34.170	349.4	407.1
	982°C	583.6	37.0	32.0	581.3	30.443	391.3	420.6
	<b>702</b> C	627.0	25.3	29.7	627.0	24.417	412.4	455.8
				31.3	301.7	29.034	198.9	215.1
	1850/36.0	302.9	33.1	27.1	312.7	27.479	201.4	220.2
	1010°C	314.4	32.0	27.1	312.7	21,517	201.4	
20	1976/28.1	90.0	19.7	29.2	88.5	16.627	33.9	48.8
20		91.5	30.3	31.9	90.6	29.001	37.3	47.9
	1080°C	68.6	35.3	32.2	. 68.4	28.869	17.3	27.6
		00.0	33.3				43.7	-
		(INTERRUP	TED TESTS)				41.4	-
		<b>(1</b>					38.7	-
			31.4	30.8	323.9	24,403	160.1	207.7
26	2012/14.5	324.1		31.9	481.1	29.581	129.9	299.6
25	1100°C	481.4	30.9	31.1	549.2	25.622	304.4	375.5
	1100 €	551.7	29.9	31.1	344.2	EJ.UEE	256.1	-
							182.8	_
		(INTERRUP	IED TESTS) ·				101.5	_
	•						101.5	_
	2050/15.0	243.4	36.1	35.0	243.3	20.614	143.1	174.2
30	1121°C	374.8	12.1	20.3	374.7	11.743	166.6	280.4
	2100/12.0	463.6	15.4	25.9	463.3	13.594	245.7	363.3
	1149°C			25.9	487.1	19.550	25.7	118.9
	1147 C	488.0	20.3	23.9	407.1		_,,	

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The method and results of W and Re microstructural segregation investigation undertaken on fully solutioned and partially solutioned CMSX-12Ri test specimens are reported in Table 26 below. The investigation indicated that it is desirable to minimize the amount of microstructure-contained residual eutectic and that for fully solutioned specimens, the solution treatments developed for the invention are successful in minimizing elemental segregation, which is important in attaining optimized mechanical properties and microstructural stability.

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#### TABLE 26

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Attoy: CMSX-12 Ri 9.525mm

Test Specimen:/(3/8")Diameter Solid Bar

Specimen Condition:

Fully Solutioned

Solutioned with 2.0% Residual Eutectic

10

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Analyses Method: Microprobe Analyses

- Random array of 350 points across a section at right angles to the growth direction
- Seven line scans, 51° apart, 50 point analyses per line

The standard deviation of the  $\boldsymbol{\mathsf{W}}$  and  $\boldsymbol{\mathsf{Re}}$  measurements are the measure of homogeneity

#### Results:

CHSX-12 RI	Standard I	Deviations
	u	Re
Fully Solutioned 2X Residual Eutectic	0.27 0.36	0.50 0.90
Comparison		
Typical CHSX-4	0.57	0.60

25

Table 27 below reports results of burner rig hot corrosion test undertaken with the CMSX-12Ri specimen. The measurements were taken at the bar location which experienced the maximum attack, i.e., 900°C (1652°F) location, with the results showing the DS MAR M 002 alloy experienced approximately 20X more metal loss than the CMSX-12Ri specimen. Visual observation showed a similar result for the CMSX-10Ri alloy. Both CMSX-10Ri alloy and CMSX-12Ri alloy showed similar resistance to attack as CMSX-4 alloy based on visual specimen review at 60, 90 and 120 hours.

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#### TABLE 27

#### HOT CORROSION

#### **METHOD**

- Burner Rig 1742°F (950°C)
   2 ppm salt, standard fuel
- Heasurements taken at point of maximum attack which was at 1652°F (900°C)
- Heasurements reported were taken at the average minimum diameter of useful metal

#### RESULTS

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+ 90 Hour Test

Alloy	Initial Dia.	Useful Dia.	Per Side
DS Mar H 002	6.88 mm	5.14 mm	.87mm (.034")
CHSX-12Ri	6.86 mm	6.78 mm.	.04 mm (.0016#)

Table 28 below reports the results of cyclic oxidation tests undertaken at 1100°C (2012°F) with Mach 1 gas velocity. The CMSX-12Ri specimen was similarly resistant to oxidation attack at 1100°C (2012°F), however, it was not as good as CMSX-4 at approximately 1030°C (1886°F) exposure.

TABLE 28

#### Cyclic Oxidation Test

15 Minute Cycles to 2012°F (1100°C), Cooled To Ambient Between Cycles

Mach 1 Gas Velocity

89 Hours Total With 77 Hours at 2012°F

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CHSX-12 Ri

RESULT: at 1100°C

Approx. 0.1 mm loss per side for every 300 cycles

CMSX-4

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Approx. 0.1 mm loss per side for every 380 cycles

at 1030°C

CHSX-12 Ri

Approx. .105 mm loss per side after 355 cycles

CMSX-4

Approx. .03 mm toss per side after 355 cycles

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CMSX-12Ri elevated temperature tensile data is reported in Table 29 below, while the results of impact tests are reported in Table 30 below. The CMSX-12Ri elevated temperature impact strength minimum is similar to CMSX-4 and its maximum occurring at 950°C (1742°F), is better.

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TABLE 29

#### TENSILE DATA

CHSX-12 RI ALLOY

TABLE 30

Test Temp	°C	LAUE	0.1% Yld ksi	0.2% Yld ksi	UTS ksi	Elong X	RA %
1382	750	2.3*	150.0	160.8	188.7	13	14
1382	750		153.6	165.1	190.0	13	15
1562	850	6.2°	136.5	130.8	152.3	27	24
1562	850	6.2°	135.0	128.9	160.1	25	23
1742	950	5.6°	92.7	89.2	125.3	24	30
1742	950	5.6°	99.9	106.2	129.2	24	32
1922	1050	3.8°	69.5	74.3	104.1	19	36
1922	1050	3.8°	72.4	77.6	106.0	19	36

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## IMPACT DATA

CMSX-12 Ri 8.89mm (0.35 Inch)Diameter Plain Cylindrical Specimens

	Test Temperature, 'F				
	1382 750°C	1562 850°C	1742 950°C	1922 1050°C	
CMSX-12 Ri (1 only)	26 J	20 J	60 J	32 J	
CMSX-4 (Ave. of 4)	26 J	21 J	42 J	45 J	

Further Impact Property Comparison

CMSX-2 -- Min. Impact Strength 16.5 Joules

. SRR 99 -- Min. Impact Strength 20 Joules

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The results of CMSX-12Ri low cycle fatigue tests undertaken at 750°C (1382°F) and 950°C (1742°F) test conditions, with R = O, are reported in Table 31 below. The data indicates that CMSX-12Ri performance is similar to CMSX-4 at 750°C (1382°F) condition, while the alloy exhibits approximately 2.5 times the typical CMSX-4 life at 950°C (1742°F) condition.

TABLE 31

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LOV CYCLE FATIGUE

CHSX-12 Ri Alloy

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R = 0 (zero to maximum stressing)

1382°F (750°C)		1742°F (950°F)			
PEAK ksi	STRESS (HPa)	Cycles	PEAK ks i	STRESS (MPa)	Cycles
142	(980)	8686	110	(760)	4427
130	(900)	11950	99	(680)	15510
125	(860)	20810	87	(600)	37430
119	(B20)	>100000	75	(520)	92580

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- . Performance is similar to CMSX-4 at 1382°F (750°C)
- Compared to CMSX-4, at 1742°F (950°C) and in the 20000 cycle region, CMSX-12Ri exhibits 2.5 times life or 15% on strength.

25

Notched low cycle fatigue test results show the CMSX-12Ri is 2-1/2 times better than CMSX-4 out to about 30000 cycles, while at 50000 cycles and above, the alloy performance is similar to CMSX-4. The results of these tests performed at 750°C (1382°F), K<sub>i</sub> = 2.0 and R=O test condition, are reported in Table 32 below.

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#### TABLE 32

NOTCHED LOW CYCLE FATIGUE	•
CMSX-1ZRi Alloy	
$1382^{\circ} f$ (750°C), $K_t = 2.0$ , $R = 0$	
PEAK STRESS ksi (MPa)	CYCLES
113.13 (780)	4879
107.33 (740)	9784
95.72 (660)	28470
84.12 (580)	49810
81.22 (560)	
78.32 (540)	> 115,000
75.42 (520)	> 115,000

Results are 2-1/2 times better than CMSX-4 out to about 30000 cycles. 35

40 High cycle fatigue test results for the CMSX-10Ri alloy are reported in Table 33 below. For 950°C (1742°F), 100 Hz, R = O test condition, the alloy exhibited about 2-1/2 times the typical CMSX-4 lives.

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Results are similar to CMSX-4 at 50000 cycles, and above.

## TABLE 33

CHSX - 108	i Alloy	
1742*!	(950°C), 100 Hz., R = 0	
PEAK STI		(H <sup>f</sup> ) CACTE2
B1.22	(560)	15.2 x 10
92.82	(640)	3.59 x 10
	(720)	0.6 x 10

\* Lives are 2-1/2 times better than Change

25

*30* 

The CMSX-10Ri and CMSX-12Ri test data indicates that adequate hot corrosion and oxidation resistance can be achieved with extremely low alloy chromium content. Additionally, extremely good thermo-mechanical fatigue tensile and impact strengths are apparent with the superalloys of this invention.

The results of alloy specimen density measurements are reported in Table 34 below.

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#### TABLE 34

## SINGLE CRYSTAL ALLOY DENSITY DATA

	ALLOY	7p2\1u2 DEN211A	DENSITY gms/cm <sup>3</sup>
10	CHSX-10A CHSX-10B CHSX-10C CHSX-10O	.324 .324 .325 .325 .325	8.97 8.99 9.00 9.00 9.00
15	CMSX-10E CMSX-10F CMSX-10G CMSX-10Ga CMSX-10H CMSX-10I	.323 .322 .322 .324 .322 .322 .322	8.54 8.91 8.91 8.91 8.91
20	CHSX-10J CHSX-10Gb (10K) CHSX-12A CHSX-12B CHSX-12C CHSX-12Cb (12D) CHSX-10Ri CHSX-10 Ri	.329 .323 .325 .326 .326 .326	9.05 9.11 8.94 9.00 9.02 9.02 9.02 9.02

The alloys of this invention are amenable to HIP processing. Specimens HIP treated as reported in Table 35 below, showed nearly complete pore closure and absence from incipient melting.

#### TABLE 35

<b>35</b>	HIP CONDITION				
	1.	Heat Specimens in the HIP vessel to 1346°C (2455°F) at minimum Argon pressure (approximately 10.34 MPa - approximately 1500 psi) and hold for 4 hours while maintaining 2455°F/1500 psi condition.			
40	2.	While maintaining the 1346°C (2455°F) operating temperature, increase the Argon pressure over 1 hour to 137.9 MPa - 20 ksi. Soak specimens at 2455°F/20 ksi condition for 4 hours.			

While this invention has been described with respect to particular embodiments thereof, it is apparent that numerous other forms and modifications of this invention will be obvious to those skilled in the art.

#### Claims

45

1.	A nickel-based superalloy comprising the following elements in percent by weight				
	Rhenium	about 5.0-7.0			
50	Chromium	about 1.8-4.0			
	Cobalt	about 1.5-9.0			
	Tantalum	about 7.0-10.0			
	Tungsten	about 3.5-7.5			
	Aluminium	about 5.0-7.0			
<i>55</i>	Titanium	about 0.1-1.2			
	Columbium	about 0-0.5			
	Molybdenum	about 0.25-2.0			
55	Titanium Columbium	about 0.1-1.2 about 0-0.5			

Hafnium about 0-0.15
Nickel + Incidental Impurities balance
said superalloy having a phasial stability number N<sub>V3B</sub> less than about 2.10.

- A superalloy according to claim 1 wherein said superalloy has a phasial stability number N<sub>V3B</sub> less than 1.85.
  - 3. A nickel-based superalloy as claimed in claim 1 in which the following elements are present in the following proportions by weight:

Tantalum 8.0-10.0
Tungsten 3.5-6.5
Aluminium 5.3-6.5
Titanium 0.2-0.8
Columbium 0-0.3
Molybdenum 0.25-1.5

Molybdenum 0.25-1.5 Hafnium 0.02-0.05

and the superalloy has a phasial stability number N<sub>V3B</sub> less than about 1.75.

4. A superalloy according to any of claims 1 to 3 further comprising the following elements in percent by weight:

Carbon about 0-0.04
Boron about 0-0.01
Yttrium about 0-0.01
Cerium about 0-0.01
Lanthanum about 0-0.01

Lanthanum about 0-0.01

A superalloy according to any of claims 1 to 4 further comprising the following elements in percent by weight:

Manganese about 0-0.04
Silicon about 0-0.05
Zirconium about 0-0.01
Sulfur about 0-0.001
Vanadium about 0-0.10

- 6. The superalloy of any of claims 1 to 5 wherein said superalloy has a chromium content of from about 1.8 to about 3.0 percent by weight.
  - The superalloy of any of claims 1 to 6 wherein said superalloy has a rhenium content of from about 5.5 to about 6.5 percent by weight.
- 8. The superalloy of any of claims 1 to 7 wherein said superalloy has a cobalt content of from about 2.0 to about 5.0 percent by weight.
  - 9. The use of a superalloy according to any of claims 1 to 8 to cast a single crystal article.
- 10. The use of a superalloy according to any of claims 1 to 8 to cast a single crystal component for a turbine engine, preferably a gas turbine blade or gas turbine vane.
  - 11. A method of producing an article which comprises casting the article as a single crystal from an alloy according to any one of claims 1 to 8 and aging the casting at a temperature of from about 1065°C to about 1165°C for about 1 to about 20 hours.

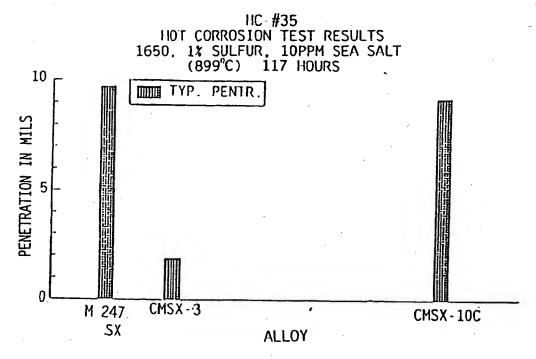
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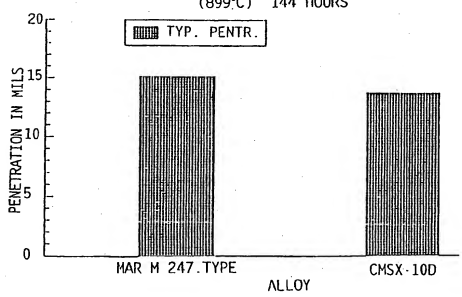
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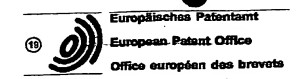
TYP. PENTR. = AVERAGE DEPTH OF CORROSION ATTACK ON PINS IN GROUP

Fig. 1

HC #35 HOT CORROSION TEST RESULTS 1650. 1% SULFUR. 10PPM SEA SALT (899°C) 144 HOURS



TYP. PENTR. = AVERAGE DEPTH OF CORROSION ATTACK ON PINS IN GROUP



1 Publication number: 0 577 316 A3

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## **EUROPEAN PATENT APPLICATION**

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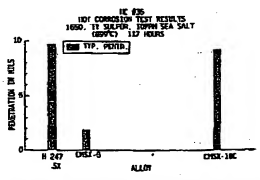
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(7) Applicant: CANNON-MUSKEGON CORPORATION 2875 Lincoln Street, P.O. Box 508 Muskegon, Michigan 49443 (US) (72) Inventor: Erickson, Gary L 874 West Street Muskegon, Michigan 49442 (US)

(4) Representative: Robinson, Anthony John Metcalf et al Kilbum & Strode 30 John Street London, WC1N 2DD (GB)

(4) Single crystal nickel-based superalloy.

A nickel-based superalicy comprises the following elements in percent by weight: from about 5.0 to about 7.0 percent rhenium, from about 1.8 to about 4.0 percent chromium, from about 1.5 to about 9.0 percent cobalt, from about 7.0 to about 7.5 percent tungsten, from about 5.0 to about 7.5 percent aluminium, from about 0.1 to about 7.0 percent itlanium, from about 0.1 to about 1.2 percent itlanium, from about 0.25 to about 2.0 percent columbium, from about 0.25 to about 2.0 percent molybdenum, from about 0.25 to about 0.15 percent hafnium, and the balance nickel + Incidental impurities, the superalicy having a phasial stability number Note 1.0 percent 2.10.



TYP. PERTR. . AVERAGE DEPTH OF CORROSION ATTACK ON PIRS IN GROUP

Fig. 1

EP 0 577 316 A3



## **EUROPEAN SEARCH REPORT**

Application Number EP 93 30 4882

ategory	DOCUMENTS CONSIDE  Citation of document with India of relevant passes	cation, where apprepriate,	Eniment to chim	CLASSIFICATION OF THE APPLICATION (backle)	
{	EP-A-0 434 996 (GENER 3 July 1991 * CLAIMS 1,3,5; TABLE		1-5,7-11	C22C19/05 C30B11/00 C30B29/52	
( _	US-A-4 849 030 (GENER 18 July 1989 * CLAIMS 16,18,19; T/		1-3,5, 7-11		
<b>\</b>	GB-A-2 235 697 (GENE) 13 March 1991 * PAGE 1, LINES 5-9;				
		_ <del></del>			
		·		TECHNICAL FEELDS SEARCHED (bel.Cl.5)	
				C22C C30B	
	A Section				
	The present morels report has b		- Sustain		
-	Place of seconds MUNICH	13 April 1994		ivalica-Bjoerk, P	
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